



# INVESTIGATION OF LITHIUM SULFUR DIOXIDE (Li/SO<sub>2</sub>) BATTERY SAFETY HAZARDS - CHEMICAL STUDIES

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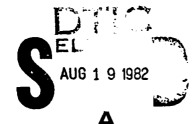
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20 ABSTRACT (Continue on feveree side if necessary and identify by block number)

The chemistry associated with the discharge and forced overdischarge of the Li/SO<sub>2</sub> cell was investigated in detail. A procedure for the quantitative determination of  $\text{Li}_2\text{S}_2\text{O}_4$  in discharged Li/SO<sub>2</sub> cells is described. The amount of  $\text{Li}_2\text{S}_2\text{O}_4$  found in cells discharged to potentials down to zero volt was in very good agreement with the discharge stoichiometry,  $2\text{Li} + 2\text{SO}_2 \rightarrow \text{Li}_2\text{S}_2\text{O}_4$ . The  $\text{Li}_2\text{S}_2\text{O}_4$  formed in the carbon cathode was also characterized by infrared, ESCA, and X-ray analyses. A number of organic com-

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pounds including CH<sub>4</sub> and 3,5-diamino-2,4-hexenenitrile have been identification forced overdischarged cells. Products identified in cells which vehicle during forced overdischarge include:  $CO_2$ ,  $CS_2$ , COS,  $H_2S$ ,  $CH_4$ ,  $C_2H_4$ ,  $C_2H_2$ ,  $Li_2S$ , and  $Li_2SO_3$ .

The performance of two types of commercial C-size Li/SO<sub>2</sub> cells (Type X and Type Z) was evaluated under abusive use conditions.

The forced overdischarge behavior of the two types of cells was markedly different. The Type Z cell displayed hazardous behavior when forced overdischarged at rates  $\geq 900$  mA. The Type X cell, on the other hand, exhibited excellent abuse resistance to forced overdischarge at currents up to 1290 mA.

The forced overdischarge of both types of cells at  $-25^{\circ}\text{C}$  was found to be a particularly hazardous operational mode. The Type Z cell was found to vent when forced overdischarged with a current of  $\geq 150$  mA. The Type X cells did not vent under similar conditions, but Li was found to plate onto the cathode. The plated Li is extremely reactive and is likely involved in exothermic reactions which produce similar reaction products as do the vented Type Z cells.

The resistance of the commercial cells to charging was also investigated briefly. Neither type of cell displayed unsafe behavior when charged after a partial discharge or a forced overdischarge.

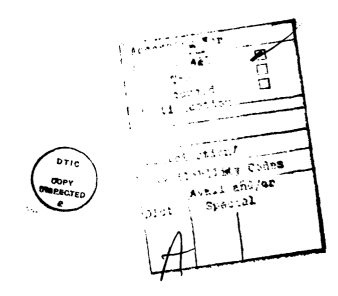
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#### **PREFACE**

A part of the experimental work in this project involves the testing of commercial  $\text{Li/SO}_2$  cells. The objective of the program is to investigate the safety hazards of the  $\text{Li/SO}_2$  system in general and not of any particular brand of commercial cell. No conclusions are presented or intended in regard to the performance of a given commercial cell.

In the present study, cells were subjected to abusive tests in order to provoke hazardous events under well defined conditions for systematic investigation. Therefore, the present data may be showing a higher frequency of hazards than would be encountered in the actual use of the cells.

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#### 1.0 INTRODUCTION

The Li/LiBr,CH<sub>3</sub>CN/SO<sub>2</sub> battery is one of the most advanced high-energy density battery systems available today. Several manufacturers currently produce Li/SO<sub>2</sub> cells in large volumes for military, industrial, and commercial applications. The outstanding features of the Li/SO<sub>2</sub> system which make it attractive to users include its high specific energy, high volumetric energy density, long shelf life, extremely stable voltage, and outstanding low temperature performance (1-10). However, the same active cell materials which provide for these desirable properties can also make the cell unsafe under certain operating conditions. As the use of the Li/SO<sub>2</sub> system increases, concern over its safety also grows.

Despite earlier claims that commercial  $\text{Li/SO}_2$  cells are sufficiently abuse resistant for specific applications (5,11,12), there has been an increasing number of reports (13) of hazardous behavior exhibited by the cell under both test and actual use conditions. Clearly, the present  $\text{Li/SO}_2$  cell has not been perfected to the point of being safe under all use conditions.

During the first phase of this program, we conducted a literature and user survey of the safety related problems of  $\text{Li/SO}_2$  cells. The detailed results of this survey have been published previously (13). The study identified three conditions under which use of the  $\text{Li/SO}_2$  system is particularly hazardous:

- (i) Forced overdischarge of Li/SO<sub>2</sub> cells. This situation, experienced by a weak cell in a seriesconnected battery, is the most frequent cause of cell or battery venting or explosion.
- (ii) Increased vulnerability of partially discharged and stored Li/SO<sub>2</sub> cells and batteries to subsequent abuse; e.g., shorts, high current pulses, overdischarge or incineration. This is a particularly hazardous condition in practical situations.
- (iii) Low temperature discharge, particularly when a cell is driven into voltage reversal and subsequently warmed up to room temperature.

The second part of this program involved an experimental investigation of the safety hazards of the  $\mathrm{Li/SO_2}$  system, particularly those associated with forced overdischarge. A major objective of this study was the thorough characterization of the cell chemistry occurring during the normal discharge, as well as the chemistry associated with hazardous use conditions. The experimental techniques and results are presented in this report.

#### 2.0 EXPERIMENTAL PROCEDURES

### 2.1 Testing and Analysis of Li/SO<sub>2</sub> Cells

The electrochemical testing, opening of unvented cells, and collection of gaseous products were carried out in a specially designed, hermetically sealed test chamber illustrated in Fig. 1. The chamber was designed to contain all compounds released from cells which vented during testing or from cells deliberately opened for analysis. An Ar or He atmosphere was maintained in the container during testing.

When placed in the test chamber, the cell was in contact with only a Teflon block on the bottom surface of the chamber and a narrow Teflon ring around 3/4 of the circumference of the cell. The safety vents were unimpeded with both type cells tested.

Cells were galvanostatically discharged using a constant current power supply with a maximum voltage of 12V. The wall temperature of the cells was measured with a copper-constantan thermocouple mechanically attached to the outside of the cell wall and calibrated against an Omega electronic ice point reference. The cell potential and temperature were measured continuously during discharge.

During the low temperature experiments, the test chamber containing the cell was placed in a Tenney environmental chamber maintained at -25°C. All cells were allowed to equilibrate at -25°C for at least 12 hours before being discharged. After the discharge or forced overdischarge, the test chamber was removed from the environmental chamber and the cells were allowed to warm to room temperature (indicated by the thermocouple mounted on the cell wall). Except where noted, the cells were allowed to stand at room temperature for at least an additional three hours before being opened.

Analyses of the cells and their contents were carried out as follows. The gases were released from the cell by piercing the end of the cell with the stainless steel plunger illustrated in Fig. 1. The plunger is moved up or down by screwing it through a series of air-tight threads in the cover of the test chamber. All volatile products released from the cell were completely retained within the test chamber.

The test chamber was equipped with two collection ports secured by valves. The gaseous compounds released into the test chamber were collected by one of two methods: i) the gases were directly transferred through one of the collection ports to evacuated gas IR cells, G. C. syringes, etc.

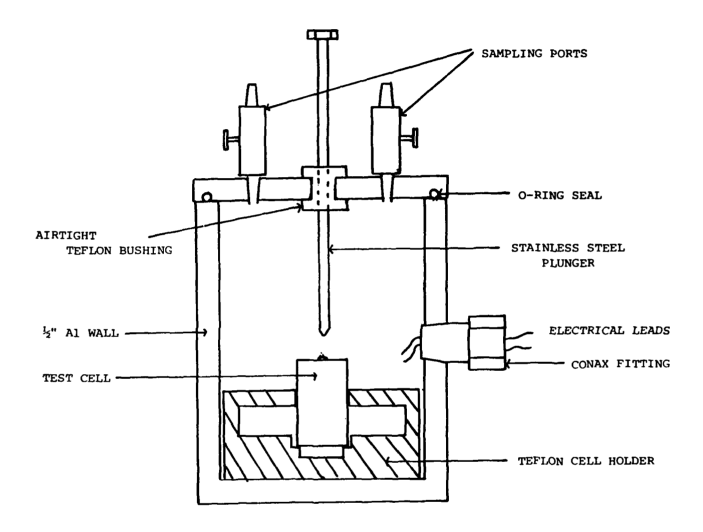


Fig. 1. Illustration of the test chamber used for the testing and opening of  $\text{Li/SO}_2$  cells.

for analysis in the respective instruments, ii) the gases were collected in an evacuated collection/storage vial and saved. The collection/storage vials had sampling ports for subsequent analysis of the gases.

The less volatile components (such as CH<sub>3</sub>CN) were collected from the cell after attaching the test chamber to a vacuum system.

A picture of the assembled test chamber and gas collection system is shown in Fig. 2. All analyses of the internal components of the cell were performed in an Ar filled glove box. Thus, throughout the entire analytical procedure, there was no atmospheric contamination. The internal cell components were removed by cutting the metal can in half, lengthwise, with a carbide grinding wheel.

Qualitative tests (14-16) were performed as follows:  ${\rm SO_3}^{-2}$  was identified by reaction with sodium nitroprusside and/or by conversion to SO<sub>2</sub> by reaction with acid;  ${\rm S_2O_4}^{-2}$  was identified by reaction with Napthol Yellow-S;  ${\rm S^{-2}}$  was identified by conversion of  ${\rm H_2S}$  after treatment with acid and/or by reaction with sodium nitroprusside in alkaline solution; CN<sup>-</sup> was identified by reaction with CuS;  ${\rm SO_4}^{-2}$  was identified by reaction with Ba<sup>+2</sup> and through elimination of other species; S was identified by melting point after extraction with CS<sub>2</sub>.

### 2.2 Instrumental Analyses

Infrared spectra were recorded on a Beckman Acculab II spectrophotometer. Solid samples were pulverized to ensure their homogeneity then pressed into KBr discs. Volatile species were analyzed with a Beckman Universal Gas cell with KBr windows. X-ray diffraction data were obtained by the Debye-Scherrer method with  $\text{CuK}_{\text{Q}}$  radiation. All X-ray samples were ground into a fine powder and sealed in quartz capillaries under an Ar atmosphere.

ESCA measurements of finely ground, discharged cathodes were made at Surface Science Laboratories, Inc., Palo Alto, CA. The samples were handled under Ar. The binding energies were referenced to hydrocarbon C(1s) at 284.6 eV. Surface elemental compositions were based on a Na<sub>2</sub>SO<sub>4</sub> reference. Mass spectra data were obtained with a Nuclide 1290G mass spectrophotometer at Biomeasure, Inc., Hopkinton, MA. Gas chromotographic analyses were performed on a Varian 920 Gas Chromotograph equipped with a thermal conductivity detector and either a 4 ft Spherocarb (Analabs) or a 4 ft Chromosorb 104 (Analabs) resin in a stainless steel column at temperatures between 25-150°C. The assignment of peak identities was based on comparison to the retention times of standard samples.

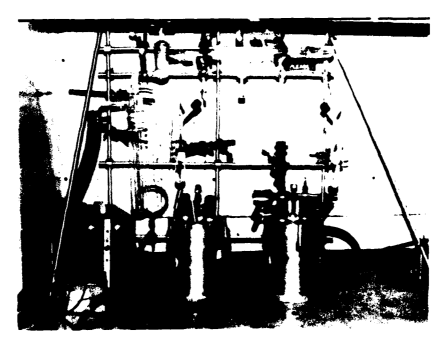




Fig. 2. Pictures of the assembled  $\text{Li/SO}_2$  test chamber and of the gas collection system.

#### 3.0 CHARACTERISTICS OF THE COMMERCIAL Li/SO2 CELLS

The evaluation of the safety hazards of  $\text{Li/SO}_2$  cells was performed with commercial C-size cells purchased from two different manufacturers. The cells are designated Type X and Type Z in this report.

The Type X cell has a rated capacity of 3.0 Ah at a discharge current of 100 mA at  $25^{\circ}$ C. The capacity of the Type Z cell is listed at 3.3 Ah at a discharge current of 135 mA.

The analysis and description of fresh Type X and Type 2 cells are summarized in Table 1.

There are a number of significant differences between the cell designs which are discussed below.

The major difference is in the cathode design. Figure 3 shows a picture of the inner section of the cathodes from the two types of cells.

The cathode of the Type X cell has a geometric area of  $\sim 178~\rm cm^2$  and was constructed with carbon pasted onto both sides of an Al grid. As seen in Fig. 3, the Al grid is attached to an Al tab at the inner end of the wrapped cathode (i.e., the center of the cell) which in turn is connected to the positive lead in the center of the cover assembly.

The cathode of the Type Z cell has a geometric area of  $125 \text{ cm}^2$  and has carbon pasted onto only one side of an Al grid. The Al grid is attached to an Al tab in the center of the cell as in the Type X cell.

Both types of cells employ a polypropylene separator. However, they apparently have different pore sizes. The material in Type Z cell is apparently more open (larger pore size) than that in Type X cell.

The Li anode in the Type X cell has essentially the same width as the cathode. It is attached to the inner wall of the can apparently by physical contact only.

The Li anode in the Type Z cell is 0.7 cm wider than the cathode. The electrodes are positioned such that the Li overlaps both edges of the cathode. The outer end of the wrapped Li is attached to a copper tab which in turn is attached to the negative lead. Post-mortem analysis of Type Z cells showed that after a discharge to 2.0V, the excess Li along the edges of the electrode package is essentially unused, insuring that the Li throughout the length of the electrode remains electrochemically accessible.

TABLE 1

DESIGN PARAMETERS<sup>a</sup> OF TYPE X AND TYPE 2 Li/SO<sub>2</sub> CELLS

|        | 13                                    | Cathode                                      | 202                    | CH3CN   | Separator            | Can                                 |
|--------|---------------------------------------|--|------------------------|---------|----------------------|-------------------------------------|
| Type x | Type X 1.25 gm (4.8 Ah) 26.0 x 3.6 cm | 4.8 gmb<br>25.5 x 3.5 x 0.09 cm              | ~9.6 gm ~2.3 gm (4 Ah) | ~2.3 gm | <b>Polypropylene</b> | Ni-plated steel with vent on bottom |
| Type Z | 1.73 gm<br>(6.68 Ah)<br>26.0 x 3.2 cm | 3.0 gm <sup>b</sup><br>25.0 x 2.5 x 0.075 cm | ~8.8 gm (3.7 Ah)       | ~1.5 gm | Polypropylene        | Ni-plated steel with vent on side.  |

arme values reflect the analysis of one cell of each type.  $\label{eq:bincludes} \textbf{bincludes weight of Al grid.}$ 



Fig. 3. Photograph of the inner section of the cathodes from the Type Z (upper) and Type X (lower) cells.

In contrast, after a discharge to 2.0V the Li remaining in the Type X cell is very thin throughout the entire length of the electrode, making it likely that most of the Li is electrochemically inaccessible during overdischarge.

The differences in the cell designs suggest that in actual use the Type X cell is anode limited, while the Type Z cell is cathode limited. Without a reference electrode in the cell this cannot be verified; however, the discharge characteristics of the cells seem to support this.

Typical voltage profiles of the discharge and forced overdischarge of Type Z and Type X cells are shown in Figs. 4 and 5, respectively. The potential profile of the Type Z cell during forced overdischarge is characterized by small negative voltages (< -500 mV) which do not show substantial changes during continued overdischarge. This behavior is consistent with an end of cell-life caused by cathode limitations.

The forced overdischarge of the Type X cell is characterized by a deep reversal of the cell voltage. After a short period in deep reversal, the cell voltage generally returns to less negative values, but consistently displays large, rapid voltage fluctuations throughout the forced overdischarge. This type of behavior is most likely to occur because of disconnection or depletion of the Li at the end of useful cell life.

It should be emphasized that even though the two types of cells are apparently of different designs, the same forced overdischarge products have been identified from both (see later).

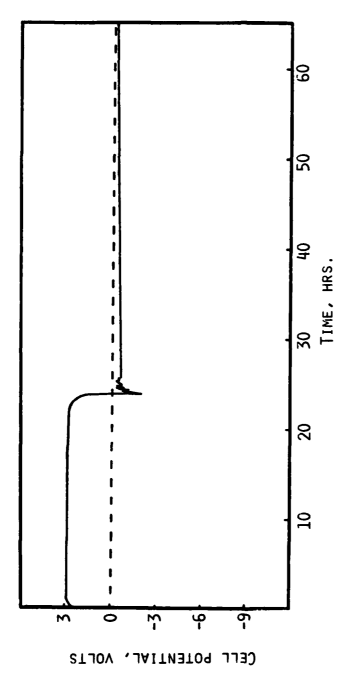
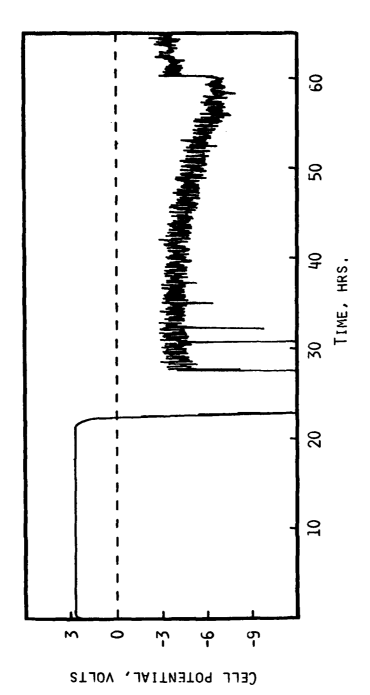


Fig. 4. The discharge and forced overdischarge of a Type 2 Li/SO<sub>2</sub> cell at 150 mA ( $\sim 1.20$  mA/cm<sup>2</sup>).



The discharge and forced overdischarge of a Type X Li/SO2 cell at 150 mA  $(\sim\!0.85~\text{mA/cm}^2)$  . Fig. 5.

### 4.0 INVESTIGATION OF THE DISCHARGE REACTION IN Li/SO2 CELLS

A series of commercial Li/SO $_2$  cells were discharged, either galvanostatically or resistively, to cell potentials down to 0V at 25 $^{\rm O}$ C. At the end of the discharges the cells were opened and analyzed. The conditions of the discharges are listed in Table 2.

- 4.1 Quantitative Analysis and Characterization of the Li/SO<sub>2</sub> Cell Discharge Product
  - 4.1.1 Quantitative Analysis of Li<sub>2</sub>S<sub>2</sub>O<sub>4</sub> in Carbon Cathodes

The discharge reaction of the Li/SO $_2$  cell results in the formation of an insoluble product in the carbon cathode. The formation of the product renders the initially flexible carbon cathode very stiff and brittle. Chemical spot tests with Napthol Yellow-S confirmed that discharged cathodes contained Li $_2$ S $_2$ O $_4$ . Li $_2$ S $_2$ O $_4$  was not detected in any other part of the cell.

Quantitative determination of the  ${\rm Li}_2{\rm S}_2{\rm O}_4$  in the cathodes of eight cells was carried out using the following procedure: The cathode was carefully removed from the cell inside a glove box, washed with acetonitrile to remove LiBr, and dried under vacuum. The entire cathode was then added to a deaerated ammoniacal silver nitrate solution (1N AgNO $_3$  in 9N NH $_4$ OH). The AgNO $_3$  was in a 50% to 100% excess over that required for the reaction of Eq. 1, based on the electrochemical capacity (to 2.0V) of each cell.

$$2Ag^{+} + S_{2}O_{4}^{-2} \rightarrow 2Ag_{(s)} + 2SO_{2(g)}$$
 (1)

The mixture was maintained under an Ar atmosphere and stirred for at least five hours. The mixture was then filtered and the solids repeatedly washed with 6N NH $_4$ OH until free of excess Ag $^+$ . The remaining solids (Ag $^{\rm O}$ , C, Al, Teflon) were added to an excess of HNO $_3$  (based on the amount of Ag $^{\rm O}$  formed in Eq. 1) and stirred overnight to dissolve the elemental silver. The mixture was then filtered and the remaining solids repeatedly washed in 6N HNO $_3$  until free of silver. The filtrates were combined and diluted to a convenient (typically 500 ml) volume with distilled water.

Aliquots (5-25 ml) of the Ag $^+$  solution were neutralized with 6N NaOH, then adjusted to a pH of 4-5 with 3N NHO $_3$ . The solutions were then boiled for five minutes to remove nitrogen oxides. After cooling to room temperature, the Ag $^+$  was determined by titration with a standard (0.1N) KSCN solution using a ferric ammonium sulfate indicator.

TABLE 2
SUMMARY OF Li/SO<sub>2</sub> CELLS DISCHARGED DOWN TO 0.0V

|                   | Current                               | Capacity | (mA)    |
|-------------------|---------------------------------------|----------|---------|
| Cell_             | (mA)                                  | to 2.0V  | to 0.0V |
| X-1               | 150                                   | 3400     | -       |
| X~2               | 160                                   | 3310     | ~       |
| X-3               | 150                                   | 3310     | -       |
| X-4               | 150                                   | 3200     | ~       |
| <b>z-1</b>        | 150                                   | 3360     | -       |
| X-5               | 750                                   | 2280     | _       |
| X-16 <sup>a</sup> | 100                                   | 1500a    | -       |
| X-19              | 150                                   | 3450     | 3560    |
| X-20              | 150                                   | 3300     | 3350    |
| X-14              | Resistive discharge across $10\Omega$ | 3140     | 3350    |
| X-22              | Resistive discharge across $10\Omega$ | 3300     | _       |

 $<sup>^{\</sup>mathrm{a}}\mathrm{Cell}$  was partially discharged.

Species which could be present in solution such as  $SO_3^{-2}$ ,  $S_2O_3^{-2}$ ,  $SO_4^{-2}$ ,  $SCN^-$ ,  $Br^-$ ,  $Al^{+3}$  were found not to interfere with the determination.

Experiments showed that  $\text{Li}_2S$  does interfere with the analysis. However, chemical spot tests with  $\text{Cu}^{+2}$ ,  $\text{Ca}^{+2}$  and  $\text{Ag}^+$  along with X-ray data (see below) gave no evidence for the presence of  $\text{Li}_2S$  in the cathode.

The results of the dithionite analyses and the experimental discharge parameters of each of the eight cells are presented in Table 3.

Three of the cells, X-1, X-2, and Z-1 were galvanostatically discharged to a cell potential of 2.0V and one cell, X-19, to a cell potential of 0.0V at 150 mA.

The amount of  $\rm Li_2S_2O_4$  found in the cells is in good agreement with the measured electrochemical capacities. The small scatter in the data is reasonable considering the relatively tedious procedure required for quantitative removal of the cathode containing the  $\rm Li_2S_2O_4$  from commercial cells.

The amount of  $\text{Li}_2\text{S}_2\text{O}_4$  found in cell X-16, which was partially discharged, is slightly high. However, it should be noted that the absolute error in the analysis (300-400 mg of  $\text{Li}_2\text{S}_2\text{O}_4$ ) is similar to that obtained in the fully discharged cells.

The amount of  $\text{Li}_2S_2O_4$  formed in cells X-17 and X-21, which were forced overdischarged, was approximately 10% greater than the electrochemical capacity to 0 volt. These results indicate that at least part of the charge passed during forced overdischarge results in the formation of  $\text{Li}_2S_2O_4$  in the cathode. The data show that further reduction of  $\text{Li}_2S_2O_4$  to other sulfuroxy species does not occur during forced overdischarge.

Cell X-25 was discharged for 1600 mAh at a current of 100 mA then recharged an equivalent amount. Analysis of the cathode found, within experimental error, no  $\rm Li_2S_2O_4$ .

Thus, the analytical results in Table 3 establish that under the conditions investigated,  $\text{Li}_2\text{S}_2\text{O}_4$  is the major and most likely the sole product formed on the cathode during discharge of  $\text{Li}/\text{SO}_2$  cells down to 0V. Furthermore, the data verify that the stoichiometry of the discharge reaction is

$$2Li + 2SO_2 \rightarrow Li_2S_2O_4 \tag{2}$$

The results from cell X-25 also verify that the cathodic reaction is reversible (see Section 7.0).

TABLE 3

DITHIONITE DETERMINATION IN COMMERCIAL C-SIZE LI/SO2 CELLS DISCHARGED AT 25°C

|                   |                              | Elect   | trochemical | Electrochemical Capacity (mAh)    |   |                  |
|-------------------|------------------------------|---------|-------------|-----------------------------------|---|------------------|
| Ce11              | Discharge<br>Current<br>(mA) | to 2.0V | to 0.0V     | Including Forced<br>Overdischarge | S <sub>2</sub> O <sub>4</sub> <sup>-2</sup> Found (mAh) | % of Theoretical |
| x-1               | 150                          | 3400    | 1           | ſ                                 | 3410  | 100.3            |
| x-2               | 150                          | 3310    | ı           | ſ                                 | 3430  | 103.6            |
| 2-1               | 150                          | 3360    | 1           | ſ                                 | 3250  | 96.7             |
| 2-19              | 150                          | 3450    | 3560        | ſ                                 | 3650  | 102.5            |
| x-16 <sup>b</sup> | 100                          | 1500b   | 1           | •                                 | 1650  | 110.0            |
| x-17c             | 150                          | 3420    | 3450        | 9750                              | 3790  | 109.8            |
| x-21c             | 150                          | 3480    | 3500        | 3930                              | 4000  | 114.3            |
| x-25d             | 100                          |         | 1           | ł                                 | 1   | 1                |
|                   |                              |         |             |                                   |   |                  |

<sup>a</sup>Based on the total measured electrochemical capacity above 0.0V.

bcell was partially discharged.

CCell was forced overdischarged.

dCell was discharged 1600 mAh then recharged 1600 mAh.

# 4.1.2 ESCA Analysis of Li<sub>2</sub>S<sub>2</sub>O<sub>4</sub>

Cell X-14 was discharged across a  $10\Omega$  resistor until the voltage fell to 0 volt. The capacity of the cell was 3350 mAh. The discharge curve is shown in Fig. 6.

The ESCA spectrum of the cathode is shown in Fig. 7. A high resolution scan of sulfur, showing the S(2p) binding energy, is given in Fig. 8.

The binding energy of dithionite S is not available in the literature. Therefore, a sample of  $\text{Na}_2\text{S}_2\text{O}_4$  (Fisher Scientific) was used as a standard.

The binding energies of the cathode containing the discharge product and  $Na_2S_2O_4$  are given in Table 4. The peak at 166.3 eV in both the  $Li_2S_2O_4$  and  $Na_2S_2O_4$  corresponds to the binding energy of S(2p) in  $S_2O_4^{-2}$ . An analysis of the binding energies of S(2p) versus the charge on S shows that the value of 166.3 eV lies where expected for S with a formal charge of +3.

The high resolution scan for S shows an additional peak at 170.4 eV in both the cathode and  $Na_2S_2O_4$  samples. The peak accounts for ~14% of the surface S in the cathode sample.

Cell X-22 was also discharged across a  $10\Omega$  resistor but only to 2.0V. The cell yielded a capacity of 3300 mAh. An ESCA analysis of the surface sulfur was identical to that of the previous cell, showing two peaks for S at 166.3 and 170.4 eV. A time dependent study showed that the intensity of the 170.4 eV peak increased relative to that of the 166.3 eV peak with increased exposure of the sample to the atmosphere. Thus the peak at 170.4 eV can be attributed to the slight decomposition of the surface dithionite due to exposure to the atmosphere during sample preparation and not to other discharge products.

There is no evidence for the presence of any other sulfur species such as  $S_2O_3^{-2}$  (160.9, 166.9 eV),  $SO_4^{-2}$  (168.0 eV),  $SO_3^{-2}$  (167.0 eV),  $S^{-2}$  (160.8 eV) or  $S^O$  (162.2 eV) in the cathode of cells discharged to 2.0 or 0.0V.

The surface elemental compositions of the cathode of cell X-14, based on a standard sample of Na $_2$ SO $_4$ , is listed in Table 5. Lacking the identity of the minor decomposition species (the 170.4 eV peak) a stoichiometric calculation of the amount of  $\rm S_2O_4^{-2}$  in the cathode is not possible. However, estimates show that  $\rm Li_2S_2O_4$  accounts for at least 90% of the surface sulfur.

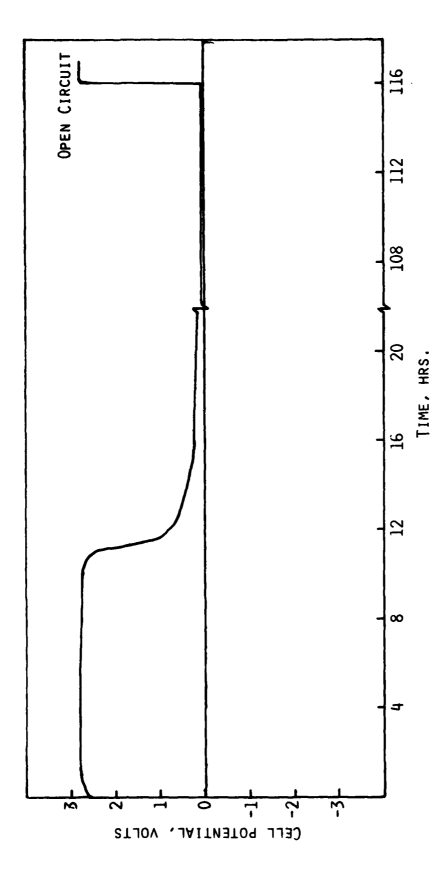


Fig. 6. Cell voltage of Li/SO<sub>2</sub> cell X-14 during discharge across a 10Ω resistor.

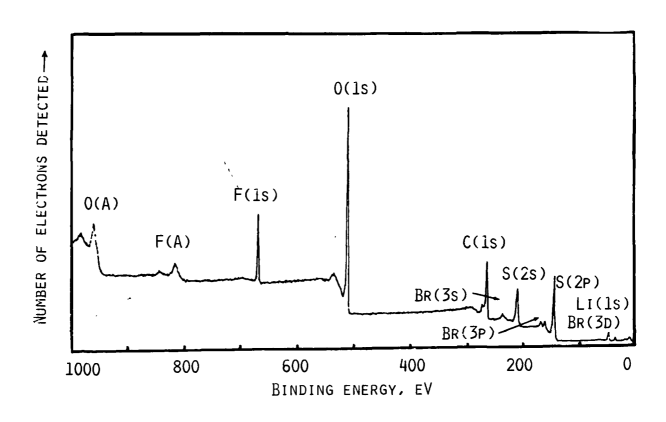


Fig. 7. ESCA spectrum of the cathode from cell X-14 after discharge to 0 volt

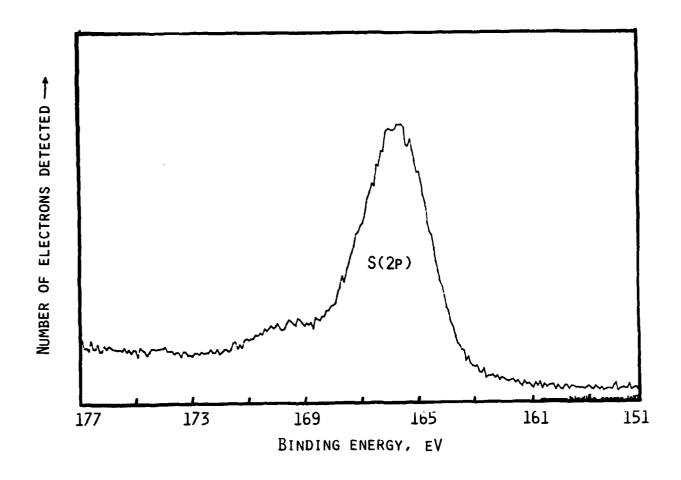


Fig. 8. High resolution ESCA scan of the surface S on the cathode from cell X-14.

TABLE 4
BINDING ENERGIES (eV) FROM HIGH-RESOLUTION SPECTRA

| Samples                               | C (ls) | 0 (ls) | S (2p)       | Li (ls) | F (1s) |
|---------------------------------------|--------|--------|--------------|---------|--------|
| Discharged<br>Cathode of<br>Cell X-14 | 284.6  | 531.7  | 166.3, 170.4 | 55.6    | 688.9  |
| Sodium<br>Dithionite                  | 284.6  | 531.8  | 166.4, 170.4 | -       | -      |

TABLE 5

SURFACE ELEMENTAL COMPOSITIONS<sup>a</sup> OF THE DISCHARGED CATHODE
FROM Li/SO<sub>2</sub> CELL X-14

| <u>c</u> | <u>o</u> | <u>s</u> | <u>Li</u> | Br  | <u>F</u> | <u>0/s</u> | Li/Sb |
|----------|----------|----------|-----------|-----|----------|------------|-------|
| 32.0     | 29.0     | 14.0     | 17.0      | 1.5 | 6.7      | 2.1        | 1.1   |

 $<sup>^{\</sup>mbox{\scriptsize a}}\mbox{\scriptsize Expressed}$  as atom percent for the detected species and based on a  $\mbox{\scriptsize Na}_2\mbox{\scriptsize SO}_4$  standard.

 $<sup>^{\</sup>mathbf{b}}\mathbf{After}$  subtracting the contribution of LiBr.

The ESCA scan in Fig. 5 also confirms the presence of all the other elements expected in the cathode. High resolution scans of C show  $\sim 2\%$  in the form of a fluorocarbon - the Teflon binder.

## 4.1.3 Infrared Characterization of Li<sub>2</sub>S<sub>2</sub>O<sub>4</sub>

The infrared spectrum of the  $\mathrm{Li}_2\mathrm{S}_2\mathrm{O}_4$  formed on the cathode of cell X-14, which was discharged to 2.0V, is shown in Fig. 9 along with the background spectrum of the Teflon bonded carbon cathode of an undischarged Type X cell (undischarged Type Z cathodes also displayed no IR absorption). The IR spectra of discharged Type Z cathodes were identical to that of the Type X.

The infrared spectra shown in Fig. 10 are from inner and outer sections of the cathode from cell X-1 (capacity 3400 mAh at 150 mA to 2.0V). The spectra are identical, showing that the same discharge product forms throughout the cathode.

The infrared spectrum of the discharge product was identical over all depths of discharge and under all discharge conditions investigated with both types of cells.

The IR spectrum of pure  $\text{Li}_2\text{S}_2\text{O}_4$  has not been reported in the literature. Therefore, the IR spectrum of the  $\text{Li}_2\text{S}_2\text{O}_4$ , formed in the carbon cathode, was compared to that of  $\text{Na}_2\text{S}_2\text{O}_4$ .

The infrared absorption frequencies of the discharge product in the cathode matrix are listed in Table 6 along with those of a sample of  $Na_2S_2O_4$  (Fisher Scientific Company). The spectrum of  $Na_2S_2O_4$  is shown in Fig. 11.

There are several similarities between the spectra of the discharge product in the cathode and  $\rm Na_2S_2O_4$ . Both spectra have a strong absorption at  $\sim\!900~\rm cm^{-1}$ . Also present in both compounds are two medium intensity absorptions at 520 and 420 cm $^{-1}$  in  $\rm Na_2S_2O_4$  and 550 and 500 cm $^{-1}$  in the discharged cathode.

The major difference between the two spectra is that the product in the cathode from the discharged cells exhibits two strong bands at 1028 and 1085 cm $^{-1}$  while Na<sub>2</sub>S<sub>2</sub>O<sub>4</sub> has only one strong band at 1050 cm $^{-1}$ . It should be noted that the frequency of this Na<sub>2</sub>S<sub>2</sub>O<sub>4</sub> band is midway between those observed in the discharged cathodes. Since the spectra were obtained on solid samples, the crystallographic effects on absorption frequencies can be significant between the Li and Na salts.

There was no evidence in any of the cathodes examined for the presence of other Li sulfuroxy compounds such as  $\text{Li}_2\text{S}_2\text{O}_3$  (1120, 1000, 670 cm<sup>-1</sup>), (970, 630, 395 cm<sup>-1</sup>), or  $\text{Li}_2\text{SO}_4$  (~1120, 645 cm<sup>-1</sup>).

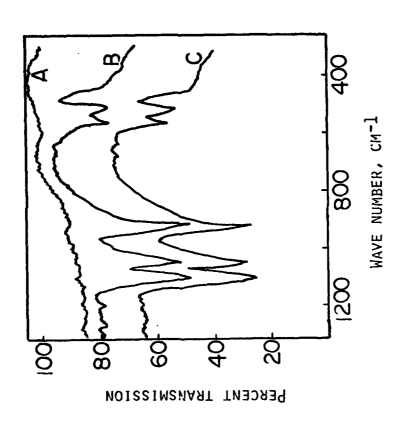


Fig. 9. Infrared spectra of Li<sub>2</sub>S<sub>2</sub>O<sub>4</sub> formed in the carbon cathode of a Li/SO<sub>2</sub> cell. Curve A: background spectrum of a fresh carbon cathode, Curve B: spectrum of a cathode after a discharge to 2.0V, and Curve C: spectrum of a cathode after a forced overdischarge.

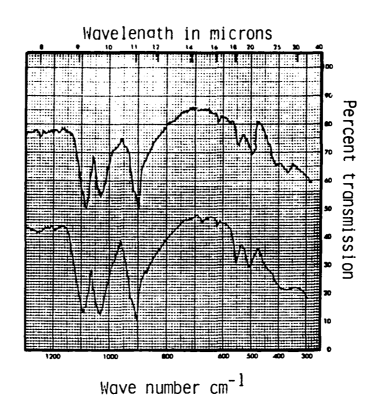


Fig. 10. IR spectra of an inner section (upper) and an outer section (lower) of the cathode from cell X-1 after discharge to 2.0V.

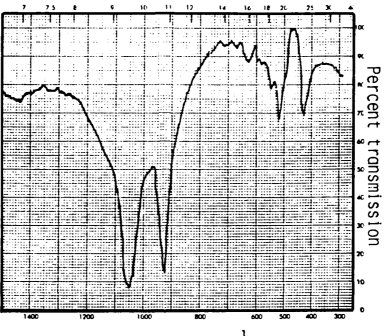
TABLE 6

MAJOR INFRARED FREQUENCIES OBSERVED IN DISCHARGED

Li/SO<sub>2</sub> CATHODES AND Na<sub>2</sub>S<sub>2</sub>O<sub>4</sub> (Fisher)

| Discharged Li/S | O <sub>2</sub> Cathodes (cm <sup>-1</sup> ) | Na <sub>2</sub> S <sub>2</sub> O <sub>4</sub> | (cm <sup>-1</sup> ) |
|-----------------|---|---|---------------------|
| 500             | (m)   | 420   | (m)                 |
| 550             | (m)   | 520   | (m)                 |
| 902             | (s)   | 551   | (w)                 |
| 1020            | (s)   | 920   | (s)                 |
| 1085            | (s)   | 1050  | (s)                 |

### Wavelength in microns



Wave number  ${\rm cm}^{-1}$ 

Fig. 11. IR spectrum of  $Na_2S_2O_4$ 

The IR spectrum of the  $\mathrm{Li}_2\mathrm{S}_2\mathrm{O}_4$  on the cathode showed no change after storage for over two months in an Ar filled dry box at room temperature. However, significant changes occurred in the infrared spectrum of the  $\mathrm{Li}_2\mathrm{S}_2\mathrm{O}_4$  (in the cathode matrix) after exposure to the atmosphere for  $\sim 16$  hours. This is seen in Fig. 12. This is consistent with the ESCA results which indicated that the  $\mathrm{Li}_2\mathrm{S}_2\mathrm{O}_4$  decomposes upon exposure to the atmosphere.

The thermal stability of the  ${\rm Li}_2{\rm S}_2{\rm O}_4$  (in the cathode matrix) was also briefly investigated. Up to  ${\sim}170^{\rm O}{\rm C}$  there is no change in the infrared spectrum of the  ${\rm Li}_2{\rm S}_2{\rm O}_4$  (again in the cathode matrix). Between  $170{\sim}200^{\rm O}{\rm C}$  the infrared spectrum shows a substantial change indicating the thermal decomposition of  ${\rm Li}_2{\rm S}_2{\rm O}_4$ . This is depicted in Fig. 13 which shows the infrared spectra of two samples of  ${\rm Li}_2{\rm S}_2{\rm O}_4$  which were heated to  $170^{\rm O}{\rm C}$  and  $200^{\rm O}{\rm C}$ , under vacuum, for one hour. This observation substantiates previous thermal studies (17-19) which assigned an exothermic transition observed in cathodes from discharged cells at  ${\sim}180^{\rm O}{\rm C}$  to the decomposition of  ${\rm Li}_2{\rm S}_2{\rm O}_4$ .

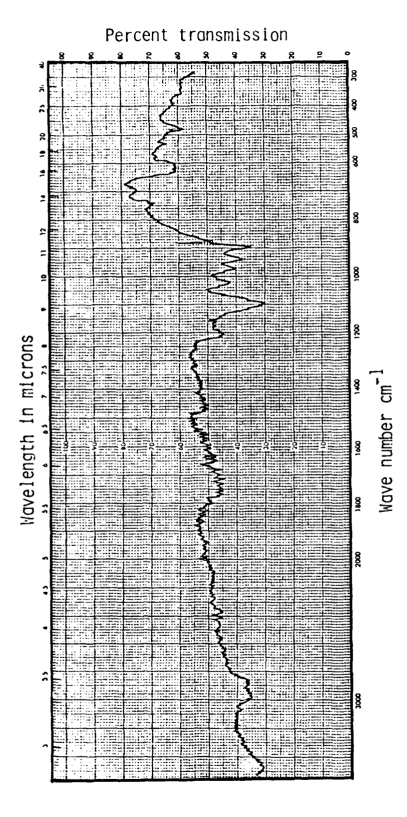
#### 4.1.4 X-Ray Analyses of Discharged Cathodes

X-ray powder diffraction data were collected on the cathodes of cells X-14, X-22, X-2, and on a predominantly white material removed from the cathode of cell X-3 by shaking it in a small amount of CH<sub>3</sub>CN. This whitish material gave a positive test for  $\rm S_2O_4^{-2}$  and had the infrared spectrum of Li<sub>2</sub>S<sub>2</sub>O<sub>4</sub>.

The four powder patterns are tabulated in Table 7. The X-ray patterns of all four cathodes are essentially identical except for variations in the relative intensities of some lines. The data clearly show that the same product is present in all the cathodes. The powder pattern also contains lines corresponding to those of LiBr (3.18, 2.75, 1.94, 1.66, 1.54, 1.38, 1.26, 1.23, 1.12 and 1.06 Å). There is no evidence in the X-ray data for the presence of Li<sub>2</sub>S, Li/Al alloy or Li/C intercalates (Li/C intercalates would probably not be detected even if present because of the amorphous nature of the carbon) in the cathodes of discharged cells.

#### 4.2 Analyses of Volatile Species in Discharged Li/SO<sub>2</sub> Cells

The gases and volatile species present in discharged  $\text{Li/SO}_2$  cells were qualitatively examined by infrared and/or gas chromatographic analyses. In all of the cells examined (except cells X-2 and Z-1) which were discharged to potentials down to 0.0V, SO<sub>2</sub> and CH<sub>3</sub>CN were the only species detected. Analyses of cells X-2 and Z-1 revealed trace amounts of CH<sub>4</sub> in addition to SO<sub>2</sub> and CH<sub>3</sub>CN.



IR spectrum of "Li $_2$ S $_2$ O $_4$ " from a discharged Li/SO $_2$  cathode (cell X-3) after exposure to the atmosphere for 16 hours.

### Wavelength in microns

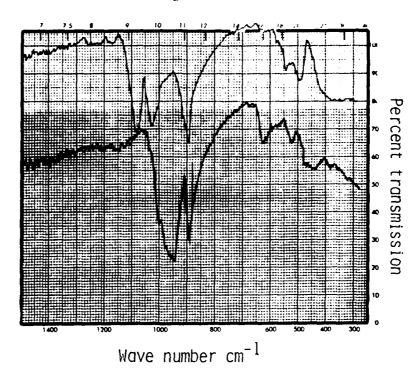


Fig. 13. IR spectra of "Li $_2$ S $_2$ O $_4$ " from the cathode of a discharged Li/SO $_2$  cell showing the effect of temperature on its stability. Top spectrum was obtained after heating to 170°C and bottom spectrum after heating to 200°C.

TABLE 7

X-RAY<sup>a</sup> DIFFRACTION DATA OF CATHODES FROM Li/SO<sub>2</sub> CELLS

|      |                  | Cell | K-22 <sup>C</sup> | Cell        | x-2 <sup>d</sup> |      | Cell X-3 <sup>e</sup> |  |  |
|------|------------------|------|-------------------|-------------|------------------|------|-----------------------|--|--|
| d, A | I/I <sub>O</sub> | d, A | I/I <sub>o</sub>  | <u>d, A</u> | I/I <sub>o</sub> | d, A | I/I <sub>o</sub>      |  |  |
| 5.10 | < 5              | 5.09 | 10                |             |                  |      |                       |  |  |
| 4.35 | 90               | 4.35 | 75                | 4.33        | 80               | 4.37 | 80                    |  |  |
| 3.73 | 80               | 3.70 | 95                | 3.69        | 90               | 3.73 | 100                   |  |  |
| 3.23 | 80               | 3.19 | 100 (br           | oad) 3.22   | 80               | 3.20 | 80                    |  |  |
| 2.96 | 50               | 2.95 | 60                | 2.93        | 40               | 2.94 | 40                    |  |  |
|      |                  | 2.74 | 40                |             |                  |      |                       |  |  |
| 2.67 | 100              | 2.66 | 100               | 2.66        | 100              | 2.70 | 100                   |  |  |
| 2.53 | 70               | 2.53 | 80                | 2.52        | 70               | 2.54 | 80                    |  |  |
| 2.43 | 5                | 2.44 | 5                 | 2.43        | 5                |      |                       |  |  |
| 2.26 | 10               | 2.25 | 30                | 2.24        | 20               | 2.25 | 15                    |  |  |
| 2.01 | < 5              |      |                   |             |                  |      |                       |  |  |
| 1.92 | 15               | 1.93 | 50                | 1.92        | 15               | 1.93 | 15                    |  |  |
| 1.80 | 10               | 1.79 | < 5               | 1.79        | 5                | 1.78 | < 5                   |  |  |
| 1.75 | 10               | 1.75 | < 5               | 1.75        | 5                |      |                       |  |  |
| 1.71 | 10               | 1.72 | 5                 | 1.72        | 5                | 1.72 | < 5                   |  |  |
|      |                  | 1.65 | 15                |             |                  |      |                       |  |  |
| 1.62 | 10               | 1.62 | 10                | 1.61        | < 5              | 1.62 | < 5                   |  |  |
|      |                  | 1.58 | 10                |             |                  |      |                       |  |  |
| 1.57 | 10               | 1.57 | 10                | 1.57        | < 5              | 1.57 | < 5                   |  |  |
| 1.53 | 10               | 1.53 | 10                |             |                  | 1.53 | < 5                   |  |  |
| 1.47 | 10               | 1.47 | 10                | 1.47        | < 5              | 1.47 | < 5                   |  |  |
| 1.42 | 10               | 1.41 | 10                |             |                  | 1.41 | < 5                   |  |  |
| 1.31 | < 5              |      |                   |             |                  |      |                       |  |  |
| 1.30 | < 5              | 1.29 | 5                 | 1.29        | < 5              |      |                       |  |  |
| 1.28 | < 5              | 1.26 | 5                 |             |                  |      |                       |  |  |
|      |                  | 1.23 | 5                 |             |                  |      |                       |  |  |
| 1.19 | < 5              | 1.19 | 5                 | 1.19        | < 5              |      |                       |  |  |
| 1.17 | < 5              | 1.16 | 5                 | 1.16        | < 5              |      |                       |  |  |
| 1.14 | < 5              | 1.14 | 5                 | 1.14        | < 5              |      |                       |  |  |
|      |                  | 1.06 | 5                 |             |                  |      |                       |  |  |
|      |                  | 0.93 | 5                 |             |                  |      | •                     |  |  |
|      |                  | 0.92 | 5                 |             |                  |      |                       |  |  |

aDebye-Sherrer method,  $\mathtt{CuK}_\alpha$  radiation.

 $<sup>^{\</sup>mbox{\scriptsize b}}\mbox{Resistively discharged to 0.0V across }10\Omega\,.$ 

<sup>&</sup>lt;sup>C</sup>Resistively discharged to 2.0V across  $10\Omega$ .

 $<sup>^{</sup>m d}$ Galvanostatically discharged at 160 mA to 2.0V.

 $<sup>^{</sup>m e}$ Galvanostatically discharged at 150 mA to 2.0V, white material separated.

#### 4.3 Li Anodes in Discharged Cells

In each of the discharged cells examined, excess, unused Li was found throughout the length of the jelly roll electrode. The Li was thickest at the point where it was attached to the negative lead in both types of cells. In the Type X cell the Li was very thin throughout. There were numerous holes through the Li, especially near its edges. In the Type Z cells the edges of the Li strip anode were essentially unused while the mid portions were very thin.

#### 4.4 Conclusions

The quantitative analysis of the  $\text{Li}_2\text{S}_2\text{O}_4$  formed on the carbon cathode of the  $\text{Li}/\text{SO}_2$  cell was accomplished by a procedure based on the reduction of  $\text{Ag}^+$  by  $\text{S}_2\text{O}_4^{-2}$ . Quantitative analyses of discharged  $\text{Li}/\text{SO}_2$  cells verified that  $\text{Li}_2\text{S}_2\text{O}_4$  is the major and most likely the sole product formed on the cathode during discharges to 0.0V. The  $\text{Li}_2\text{S}_2\text{O}_4$  was additionally characterized by infrared, X-ray, and ESCA analyses.

# 5.0 SAFETY HAZARDS DURING ROOM TEMPERATURE FORCED OVERDISCHARGE OF Li/SO<sub>2</sub> CELLS

The abuse resistance of both Type X and Type Z cells towards forced overdischarge at room temperature was evaluated. Type Z cells were tested at currents up to 1000 mA or a current density of 8 mA/cm $^2$ . Type X cells were forced overdischarged at currents up to 1290 mA, which is equal to a current density of 7.2 mA/cm $^2$ . The data show that there is a significant difference in the safety characteristics of the two types of cells during forced overdischarge.

Type Z cells were found to vent, often with flame, when forced over-discharged at currents of 900 mA or greater. At currents between 600-800 mA there was evidence of charring within the cell; however, the cells did not vent.

The Type X cells showed excellent abuse resistance. None of the cells tested vented at currents up to 1290 mA.

Table 8 summarizes the test conditions and the results of the cells tested in a forced overdischarge mode. The behavior and analyses of the cells are discussed below.

#### 5.1 Forced Overdischarge Behavior of Type X Cells

Voltage and temperature profiles of the discharge and forced overdischarge of a number of Type X cells are given in Figs. 14-18.

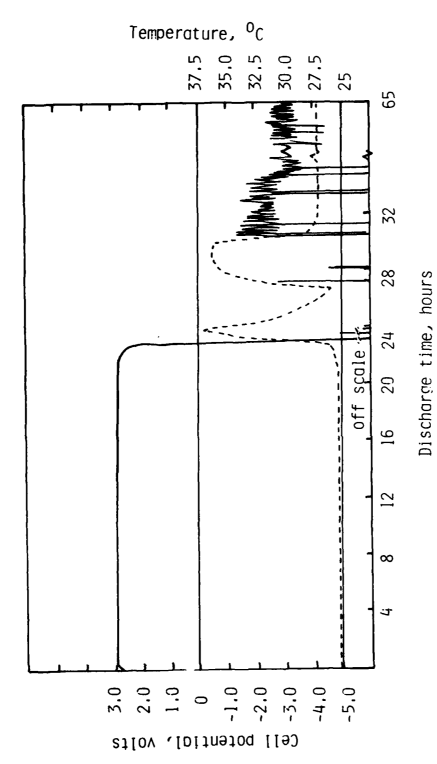
In the forced overdischarge of Type X cells at  $\leq 300$  mA, the cell potential initially fell to very negative values. When measured, the potential was approximately -12V, the voltage of the power supply. In each case, the rapid drop in cell potential at the end of discharge and the deep reversal at the beginning of forced overdischarge were mirrored by an increase in the cell wall temperature. After this point, each cell displayed slightly different behavior. In most cells, the potential rose to less negative values in 2-4 hours, but in others it remained at -12V. However, in general, further forced overdischarge resulted in large fluctuations in the cell potential which were often accompanied by changes in the cell wall temperature.

At discharge rates of  $\geq 1000$  mA the behavior of the Type X cells was slightly different. The main difference is that the cells did not experience an immediate deep reversal upon forced overdischarge. This suggest that at high rates the cells become cathode limited. However, even under these conditions none of the cells tested vented.

TABLE 8

SUMMARY OF THE RESULTS OF FORCED OVERDISCHARGE STUDIES OF LI/SO2 CELLS

| Results                                     | Did not vent. | Vented with flame. | Vented. | Vented. | Vented with flame. | Did not vent, evidence of burning within cell. | Did not vent. |
|---|---------------|--------------------|---------|---------|--------------------|--|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|---------------|
| Extent of Forced<br>Overdischarge<br>(mAh)  | 2600          | 2565               | 3600    | 340     | 3370               | 18,560   | 45,000        | 93,000        | 157,900       | 4970          | 1840          | 6320 E        | 7760          | 101,880       | 13,600        | 9750          | 3936          |
| (mAh) to                                    | 3580          | 1735               | 2000    | 2360    | 2300               | 1760   | 2100          | 3300          | 2900          | 3170          | 3390          | 3500          | 3243          | 3117          | 2525          | 3450          | 3495          |
| Capacity (mAh) to 2.0V 0.0V                 | 3540          | 1100               | 1175    | 1610    | 1530               | 1240   | 1560          | 3300          | 2840          | 3140          | 3360          | 3490          | 2700          | 2620          | 1700          | 3420          | 3480          |
| Forced-<br>Overdischarge<br>Current<br>(mA) | 150           | 1000               | 1000    | 1000    | 006                | 800  | 009           | 1000          | 750           | 300           | 300           | 150           | 1000          | 1000          | 1290          | 150           | 150           |
| Discharge<br>Current<br>(mA)                | 150           | 1000               | 1000    | 1000    | 006                | 800  | 009           | 150           | 300           | 300           | 300           | 150           | 1000          | 1000          | 1290          | 150           | 150           |
| Ce11  | 2-2           | 2-3                | 7-2     | 2~2     | 9-2                | 2-7  | 8-2           | 6-2           | z-10          | 9-X           | X-7           | 8-X           | X-11          | X-12          | X-13          | x-17          | X-21          |



Voltage (-----) and temperature (-----) profiles of the discharge and forced overdischarge of Li/SO $_2$  cell X-8 at 150 mA. Fig. 14.

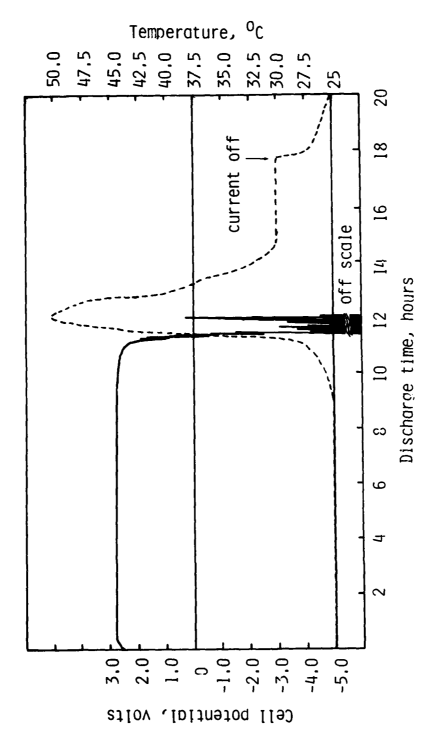
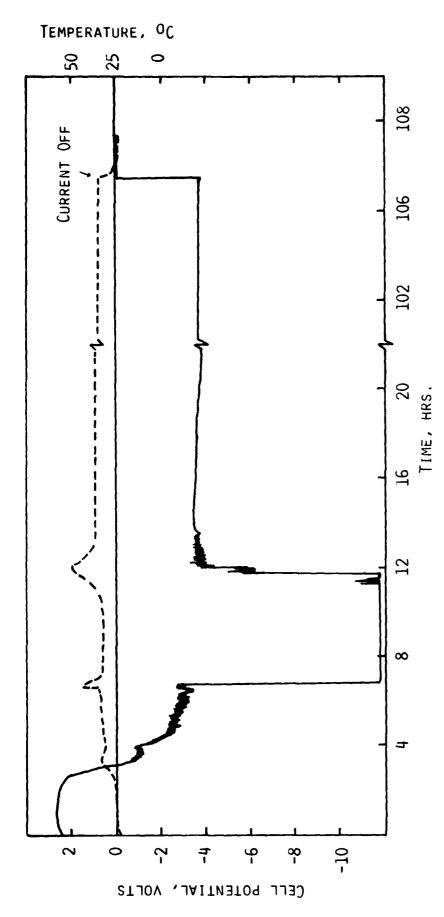
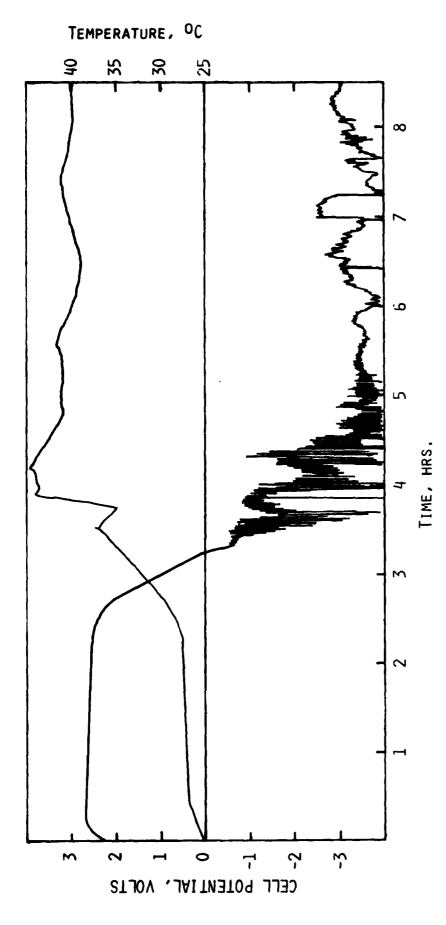


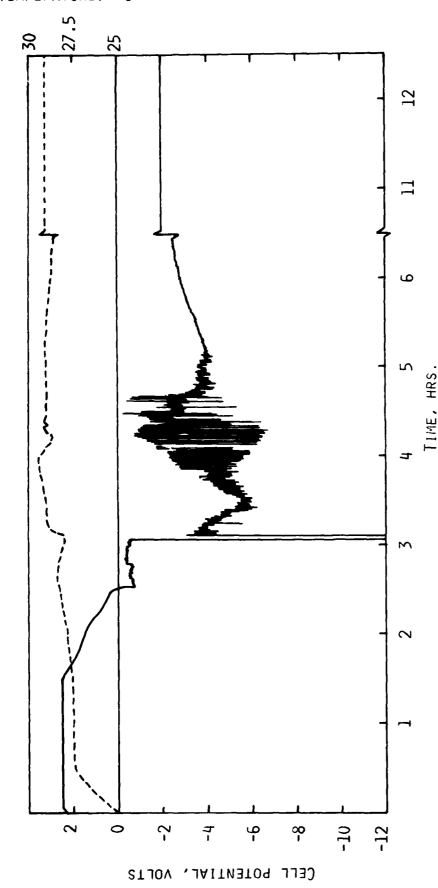
Fig. 15. Voltage (-----) and temperature (-----) profiles of the discharge and forced overdischarge of  $\text{Li/SO}_2$  cell X-7 at 300 mA.



Voltage (\_\_\_) and temperature (---) profiles of the discharge and forced overdischarge of Li/SO $_2$  cell X-12 at 1000 mA.



Voltage (\_\_\_) and temperature (---) profiles of the discharge and forced overdischarge of Li/SO2 cell X-ll at 1000 mA.



Voltage ( $\longrightarrow$ ) and temperature (---) profiles of the discharge and forced overdischarge of cell X-13 at 1290 mA.

#### 5.2 Forced Overdischarge Behavior of Type 2 Cells

The behavior of the Type Z cell on forced overdischarge at  $\leq 300$  mA was quite different as seen in Fig. 19. Once the cell potential reached 2.0V the cell rapidly went into shallow reversal. The accompanying temperature increase was less than in the Type X cells. After this point the cell potential gradually increased, levelling off usually between 0 and 0.5V, while the temperature slowly declined to room temperature. It should also be noted that the large voltage fluctuation observed with the Type X cells are not present with the Type Z cell.

At discharge rates  $\geq 600$  mA the discharge and forced overdischarge behavior was different. Figures 20-25 show the voltage and wall temperature profiles of cells forced overdischarged at rates of 600, 800, 900 and 1000 mA. In contrast to the Type X cells, the useful capacity of the Type Z cells decreased markedly with increasing discharge rates, as seen in Table 8. All the Type Z cells discharged at high rates displayed a sloping potential between 2.0 and 0 volts and each had a significant capacity near 0 volt. With continued forced overdischarge all the cells experienced large increases in the reversal potentials. All Type Z cells discharged at rates of  $\geq 900$  mA vented during forced overdischarge.

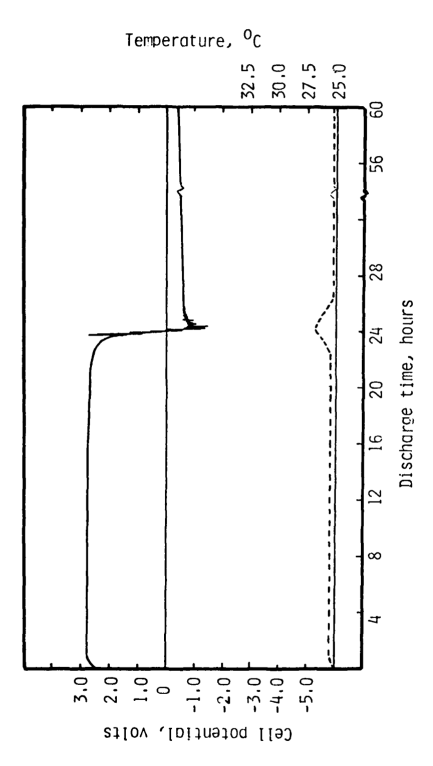
#### 5.3 Post Test Analyses of Forced Overdischarged Cells

# 5.3.1 Type X and Type Z Cells Forced Overdischarged at Rates of $\leq 300~\text{mA}$

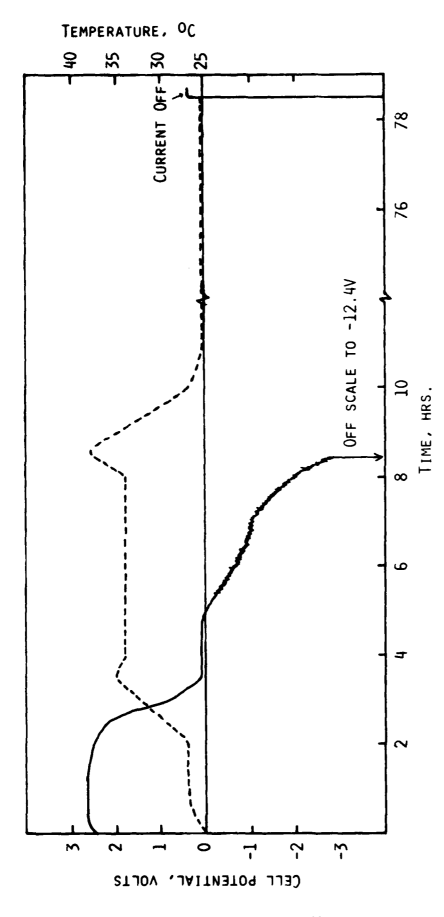
Four Li/SO<sub>2</sub> cells (X-6, X-7, X-8 and Z-2) which were forced over-discharged at low rates (i.e.,  $\leq 300$  mA) were opened and the products analyzed. All of the cells were opened within one day after the forced over-discharge was terminated except cell X-6 which was stored at room temperature for 18 days following the forced overdischarge.

Figure 26 shows the infrared spectrum of the gases released into the test chamber from cell X-7 when its top was pierced. The infrared spectrum shows the presence of  $SO_2$  -- only its most intense peak at 1361 cm<sup>-1</sup> is observed. The presence of  $CH_4$  is also confirmed by the two sharp absorptions at 3020 and 1310 cm<sup>-1</sup>. The infrared spectrum also shows the presence of another component(s) with a strong absorption at 1740 cm<sup>-1</sup>, a triplet centered at 1220 cm<sup>-1</sup> and a weak absorption at  $\sim$ 2950 cm<sup>-1</sup>.

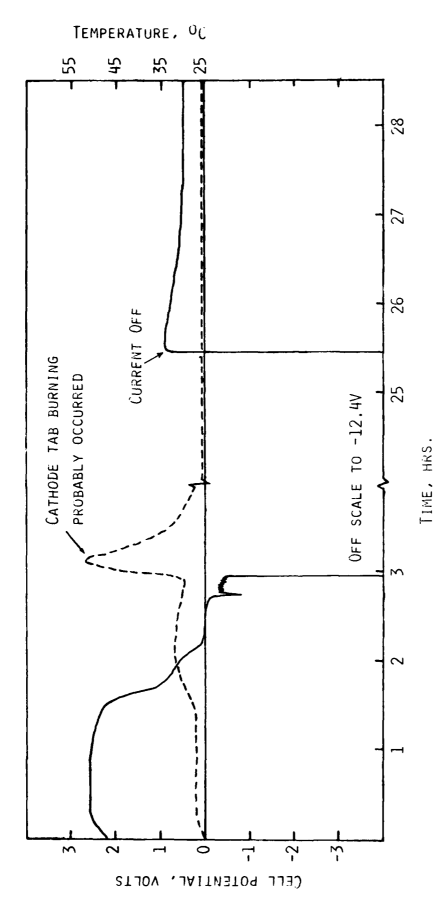
A GC analysis of the gaseous mixture showed only  $SO_2$  and  $CH_4$ . The identity of the third component observed in the infrared spectrum is not certain. The absorption at  $1740~\rm cm^{-1}$  is characteristic of the C=O stretch of an aldehyde or ketone, the weak peaks at  $\sim 2950~\rm cm^{-1}$  are in the C-H region of an aldehyde, and the triplet at  $1220~\rm cm^{-1}$  is in the region of the C-O stretch of dimerized or trimerized aldehydes. Based on the infrared data, particularly the strong absorption at  $1740~\rm cm^{-1}$ , this third component appears to be a volatile aldehyde.



Voltage (-----) and temperature (-----) profiles of the discharge and forced overdischarge of Li/SO<sub>2</sub> cell Z-2 at 150 mA. Fig. 19.



Voltage ( $\longrightarrow$ ) and temperature (---) profiles of the discharge and forced overdischarge of Li/SO<sub>2</sub> cell 2-8 at 600 mA. Fig. 20.



Voitage (——) and temperature (---) profiles of the discharge and forced overdischarge of Li/SO2 cell 2-7 at 800  $\rm mA$ . Fig. 21.

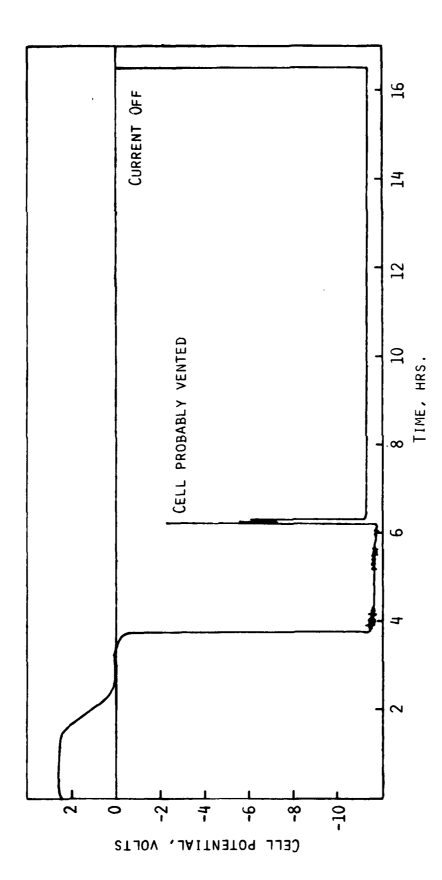
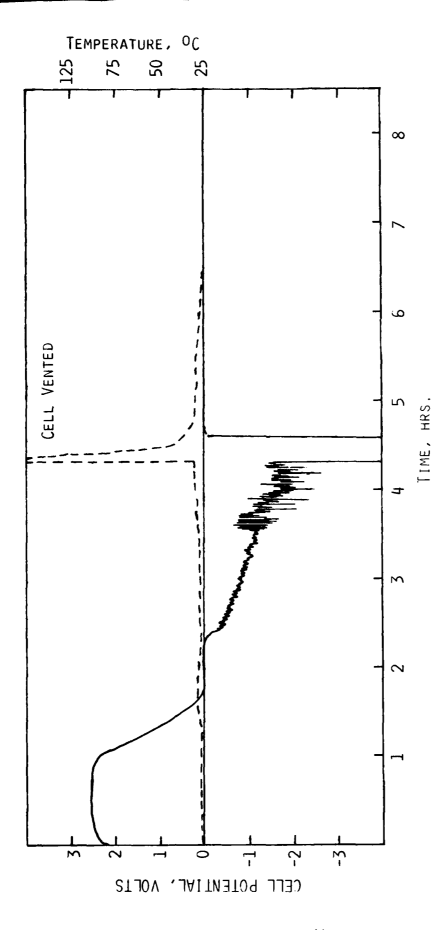
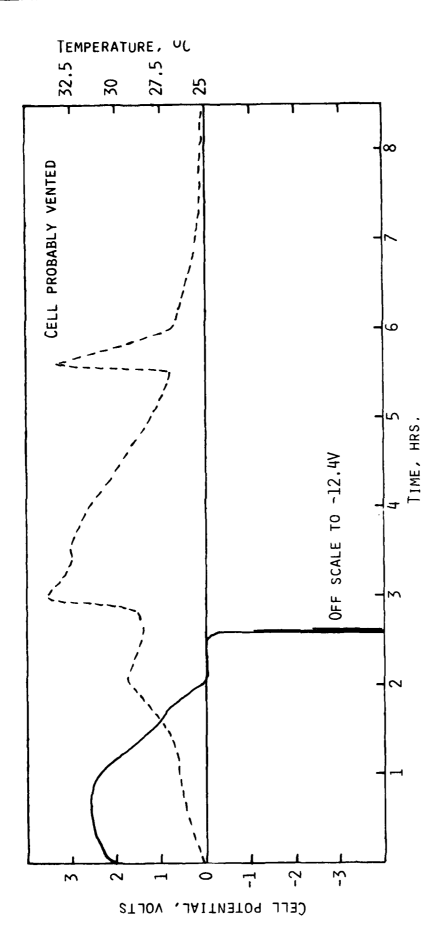


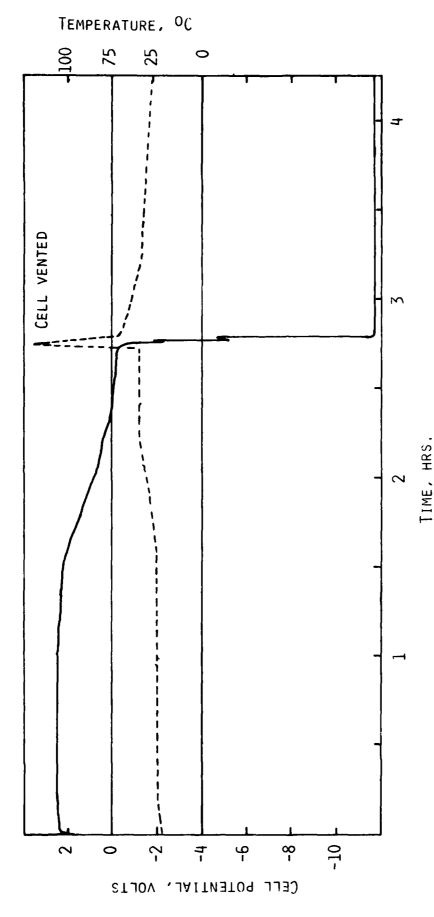
Fig. 22. Voltage profile of the discharge and forced overdischarge of Li/SO2 cell 2-6 at 900 mA.



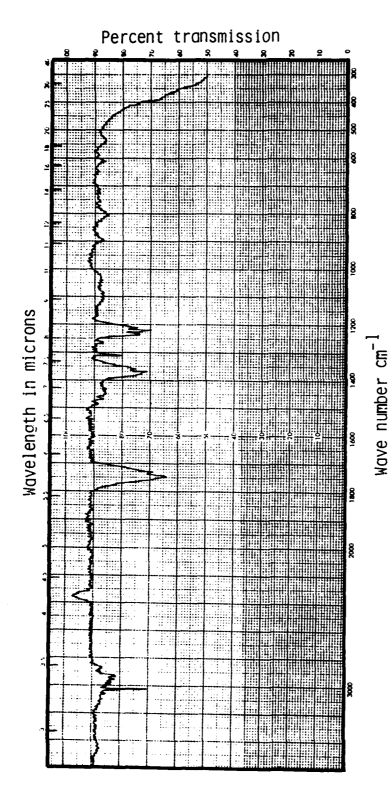
Voltage (\_\_\_) and temperature (---) profiles of the discharge and forced overdischarge of Li/S02 cell 2-3 at  $1000~{\rm mA}$ . Fig. 23.



Voltage (\_\_\_) and temperature (---) profiles of the discharge and forced overdischarge of Li/SO2 cell Z-4 at 1000 mA. Fig. 24.



Voltage (——) and temperature (---) profiles of the discharge and forced overdischarge of  $\rm Li/SO_2$  cell Z-5 at 1000 mA. Fig. 25.



ig. 26. IR spectrum of the gases from Li/SO2 cell X-7 after reversal.

The remainder of the cell was disassembled in a dry box. A small amount of Li was found on the cell wall where the Li anode was originally attached. A yellow-brown material was located throughout the rest of the anode compartment. Approximately 500 mg of the material was removed from the cell. A more detailed analysis of this material is presented later.

The infrared spectrum of the cathode from cell X-7 is shown in Fig. 27. The infrared spectrum of this cathode, even after the cell was forced overdischarged into voltage reversal, is identical to that of  $\rm Li_2S_2O_4$ . This shows that during the forced overdischarge of a  $\rm Li/SO_2$  cell no other, at least infrared detectable, species are formed on the carbon cathode. There is also no evidence for the further reduction of the  $\rm Li_2S_2O_4$  formed during discharge.

Cell X-8 was opened immediately after the current was turned off. The gases released from the cell into the test chamber were identified by IR and GC analyses as  $\rm SO_2$  and the "aldehyde" identified in the previous cell. No CH4 was detected in the gases released from the cell. The analysis of the remainder of the cell was the same as that of cell X-7.

Cell Z-2 was analyzed immediately after being forced overdischarged at 150 mA for 160% of its capacity to 2.0V. CH<sub>4</sub> and a small amount of  $SO_2$  were the only gases identified in the cell. A yellow-brown material was also found in the anode compartment of cell Z-2 and identified as the same product as found in cells X-7 and X-8. The infrared spectrum of the cathode showed only  $\text{Li}_2S_2O_4$ .

Cell X-6 was stored for 18 days before it was opened and analyzed.  $\rm SO_2$  was the only gas identified. The infrared spectrum of the cathode again showed only  $\rm Li_2S_2O_4$ . Unlike the three previous cells no yellow-brown product was found in the anode compartment. Acetonitrile was found in all the cells.

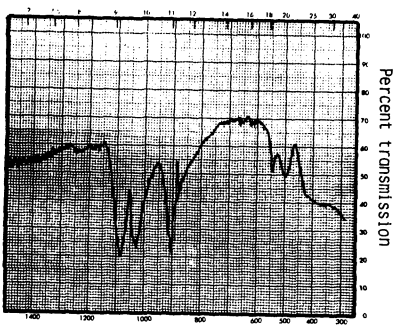
#### 5.3.2 Type X Cells Forced Overdischarged at High Rates

Three Type X cells, X-11, X-12 and X-13, were opened and examined after being forced overdischarged at rates of 1000 and 1290 mA as indicated in Table 8.

 ${\rm SO}_2$  and CH<sub>4</sub> were the only gases detected in the cells. Examination of the inside of the cells showed very little Li left in the anode compartment. In each cell there was also the brown crystal typically found in the anode of forced overdischarge cells.

Infrared analyses of the cathodes showed only the presence of  $\text{Li}_2\text{S}_2\text{O}_4$ . The Al cathode support was very brittle around the outside of the jelly roll. The Al grid also was covered with a brown compound in a few areas, however, it had formed in amounts too small to identify.

## Wavelength in microns



Wave number  $\mbox{cm}^{-1}$ 

Fig. 27. IR spectrum of the cathode of Li/SO<sub>2</sub> cell X-7 after forced overdischarge.

The separator showed an orange discoloration around the outer wind of the jelly roll electrode. Attempts to obtain spectral data on the orange material were unsuccessful.

#### 5.3.3 Type Z Cells Forced Overdischarged at High Rates

The high rate overdischarge of the Type Z cells was found to reproducibly result in cell venting, in some cases with flame. These cells were examined in greater detail.

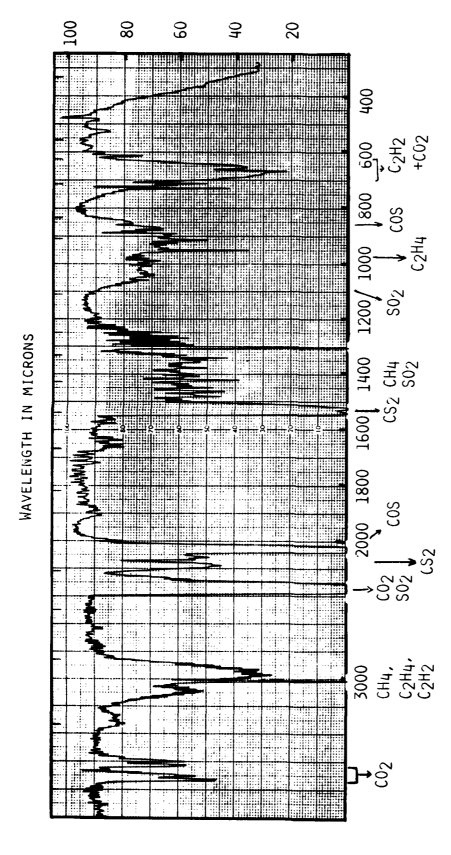
Three cells 2-3, Z-4 and Z-5 were forced overdischarged with a current of 1000 mA and Z-6 at 900 mA. All four cells vented; however, only cells Z-3 and Z-6 vented with flame.

The gases released from cell Z-3 were collected within 10 minutes after the venting. The IR spectrum of the gaseous mixture is shown in Fig. 28. Infrared and gas chromatographic analyses identified the gases as a mixture of  $CO_2$ ,  $SO_2$ ,  $CH_4$ , COS,  $CS_2$ ,  $C_2H_4$ ,  $C_2H_2$  and  $H_2S$  (identified by GConly). The entire outside of the cell was completely charred. A picture of the cell after removal from the test chamber is shown in Fig. 29. The Teflon cell holder in the test chamber was melted indicating the intense heat generated when the cell vented. A yellowish-brown material was expelled during venting. Its infrared spectrum is shown in Fig. 30. Spot tests confirmed the presence of sulfur,  $S^{-2}$ ,  $SO_3^{-2}$ , and organic compounds. The inside of the cell was fused into a solid mass as seen in Fig. 29. Thus the cathode could not be unrolled. There was no evidence of the separator or any Li left in the fused cathode. The Al tab which connects the cathode to the positive lead was completely burned away. Decomposition of the Li<sub>2</sub>S<sub>2</sub>O<sub>4</sub> on the cathode was apparent by its infrared spectrum, shown in Fig. 31, and chemical spot test. The analysis of cell 2-6 showed identical results.

The gaseous mixture from cell Z-4 was identified as containing the same species as found in cell Z-3. However, the relative amounts of  $CO_2$ ,  $CH_4$ ,  $H_2S$ ,  $C_2H_2$ ,  $C_2H_4$ ,  $CS_2$  and COS were different from those in cell Z-3. Examination of the interior of the cell showed that the separator was charred and fused to the cathode along the top edge of the jelly roll. In the core of the cell the excess separator was completely fused and again the Al tab connecting the cathode to the positive lead was completely burned away. The remainder of the cell showed no visible evidence of damage. An infrared spectrum of the cathode did not show any evidence for the decomposition of  $\text{Li}_2S_2O_4$ . Examination of cell Z-4 showed that the most intense heating occurred in the center of the cell where the Al tab connects the cathode to the positive lead.

Cell Z-5 was discharged in an inverted position and the exterior wrappings were removed before testing. Analyses showed identical results to those of cell Z-4. The Al tab on the end of the cathode was again

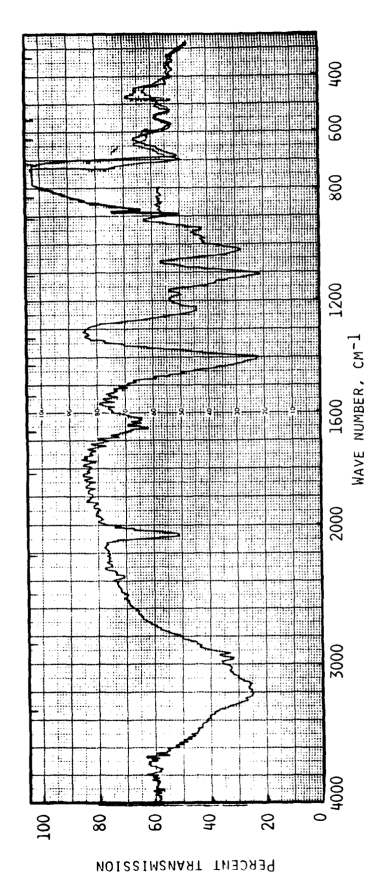
## PERCENT TRANSMISSION



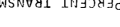
Infrared spectrum of the gases released from Li/SO $_2$  cell 2--3 after venting during 1000 mA discharge.



Fig. 29. Top photograph shows the condition of the outside of  $\text{Li/SO}_2$  cell 2-3 after venting with flame. Bottom photograph shows the completely burned interior of the same cell.



Infrared spectrum of the solid material expelled from cell 2-3 during venting.



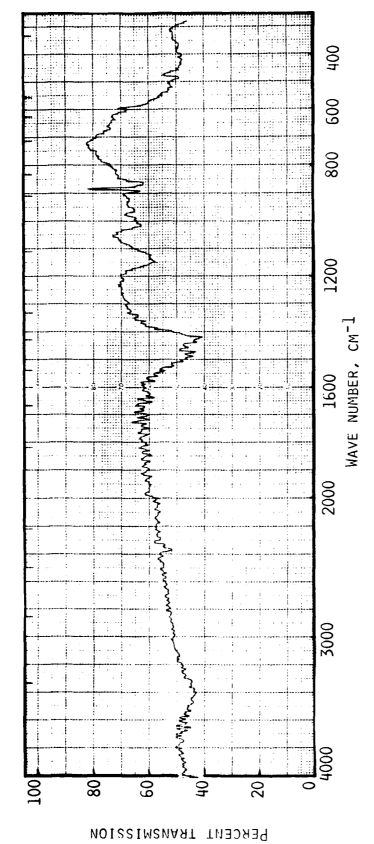


Fig. 31. Infrared spectrum of the cathode of  $\mathrm{Li}/\mathrm{SO}_2$  cell 2-3 after venting.

burned away and the separator along the top of the jelly roll was fused to the cathode.

The gaseous products found in cell Z-5 were the same as in cells Z-3 and Z-4. This indicates that the gases are formed from reactions within the cell and not from the burning of the exterior wrappings.

The fact that the area surrounding the cathode tab in the top center of the cell sustained the most intense heat in all cells, even though they had been mounted in opposite orientations, shows that the process leading to the venting is not a vapor phase reaction.

Cells Z-7 and Z-8 were forced overdischarged at 800 and 600 mA, respectively. The voltage profiles of the cells are shown in Figs. 20 and 21 and the capacities are listed in Table 8. Neither cell vented. However, post test examination revealed the cathode had become disconnected in both cells.

 ${
m CH_4}$  and  ${
m SO_2}$  were detected in cell Z-7 while only  ${
m CH_4}$  was found in cell Z-8.

A picture of the disassembled cell 2-7 is shown in Fig. 32. Examination of the cell showed that the Al tab on the cathode was burned off near the positive lead connector. The separator in the immediate area around the burned tab was also charred as seen in the picture. There were no other visible damages. The excess Li remained intact through the length of the anode compartment.

The X-ray diffraction pattern of the inner section of the cathode from cell Z-8 is tabulated in Table 9. The X-ray pattern contains lines which can be assigned to Li/Al alloy of the composition LiAl along with the lines of Li<sub>2</sub>S<sub>2</sub>O<sub>4</sub>, LiBr and Al. If the entire Al grid was converted to LiAl, the w/o of LiAl would account for  $\sim$ 7% of the total weight of the discharged cathode. In the cathode of cell Z-8, whose X-ray pattern shows the presence of unalloyed Al, the actual amount of LiAl is much less. Thus any diffraction lines of LiAl are expected to be weak, as is found in Table 9. There is no evidence in the X-ray data for the presence of unalloyed Li (2.48 Å) or any other products such as Li/C intercalates, Li<sub>2</sub>S, etc. which could form during forced overdischarge.

Cells 2-9 and Z-10 (Table 8) were discharged to 0 volt at low rates (i.e., 150 and 300 mA, respectively) then forced overdischarged at 1000 and 750 mA, respectively. When the current was increased the potential of both cells immediately fell to -12V and remained there. Neither cell vented. Post test examination of the cells showed that the cathode remained connected in both cells. However, the Al cathode grid in cell Z-9 appeared to have fallen apart over the inner section of the cathode as was seen in cell Z-8.



Fig. 32. Photograph of  $\rm Li/SO_2$  cell X-7 showing the burned Altab and separator in the top interior of the cell. The cell did not vent.

TABLE 9 X-RAY a DIFFRACTION DATA OF THE CATHODE OF CELL Z-8

| Cell Z-8    |                  | Lil  | <u> </u>         | Al   |                        |  |
|-------------|------------------|------|------------------|------|------------------------|--|
| <u>d, Å</u> | I/I <sub>O</sub> | d, A | I/I <sub>O</sub> | d, A | <u>I/I<sub>O</sub></u> |  |
| 5.09        | 5                |      |                  |      |                        |  |
| 4.37        | 40               |      |                  |      |                        |  |
| 3.70        | 90               | 3.65 | 75               |      |                        |  |
| 3.22        | 80               |      |                  |      |                        |  |
| 2.96        | 30               |      |                  |      |                        |  |
| 2.73        | 10               |      |                  |      |                        |  |
| 2.66        | 100              |      |                  |      |                        |  |
| 2.53        | 30               | •    |                  |      |                        |  |
| 2.43        | 5                |      |                  |      |                        |  |
| 2.34        | 35               |      |                  | 2.34 | 100                    |  |
| 2.24        | 30               | 2.26 | 100              |      |                        |  |
| 2.02        | 20               |      |                  | 2.02 | 100                    |  |
| 1.92        | 25               | 1.92 | 75               |      |                        |  |
| 1.79        | 5                |      |                  |      |                        |  |
| 1.70        | 5                |      |                  |      |                        |  |
| 1.65        | 10               |      |                  |      |                        |  |
| 1.62        | 10               |      |                  |      |                        |  |
| 1.59        | 10               | 1.58 | 60               |      |                        |  |
| 1.57        | 10               |      |                  |      |                        |  |
| 1.53        | 10               |      |                  |      |                        |  |
| 1.46        | 10               | 1.46 | 60               | 1.47 | 22                     |  |
| 1.43        | 15               |      |                  |      |                        |  |
| 1.41        | 10               |      |                  |      |                        |  |
| 1.29        | 15               | 1.30 | 100              |      |                        |  |
| 1.28        |                  |      |                  |      |                        |  |
| 1.26        | 5                |      |                  |      |                        |  |
| 1.22        | 19               | 1.22 | 75               | 1.22 | 24                     |  |
| 1.17        | 5                |      |                  |      |                        |  |
| 1.14        | < 5              |      |                  |      |                        |  |
| 1.12        | 5                | 1.12 | 75               |      |                        |  |
| 1.08        | 5                | 1.07 | 75               |      |                        |  |
| 1.06        | 5                | 1.01 | 85               |      |                        |  |
| 0.93        | 5                | 0.92 | 50               |      |                        |  |
| 0.91        | 5                | 0.89 | 75               |      |                        |  |
| 0.85        | 5                | 0.85 | 125              |      |                        |  |
| 0.83        | 5                | 0.83 | 100              |      |                        |  |
|             |                  | 0.80 | 60               |      |                        |  |

 $<sup>^{</sup>a}\text{Debye-Sherrer}$  method,  $\text{CuK}_{\alpha}$  radiation.  $^{b}\text{Cell}$  was discharged and forced overdischarged at 600 mA.

#### 5.4 Analysis of the Products Formed at the Anode During Reversal

In both the Type X and Type Z which did not vent when forced over-discharged (except X-6), a few hundred milligrams of a yellow-brown material was found throughout the anode compartment. Little or no Li remained in the anode compartments. The infrared spectra of the material from each cell were identical. The infrared spectrum of the material, shown in Fig. 33, indicates the presence of N-H groups (3000-3500 cm $^{-1}$ ), -C=N in a conjugated, unsaturated system (2200-2250 cm $^{-1}$ ), C=C bonds (1620-1680 cm $^{-1}$ ) and S-O bonds (900-1000 cm $^{-1}$ ).

The material used in the following analysis was obtained from cell X-8. An elemental analysis (Galbraith Labs, Knoxville, TN) gave the following composition: C, 28.22%; N, 16.63%; H, 4.33%; S, 9.10%; Br, 4.52%; and Li, 9.30%. The remainder of the sample is presumably oxygen which by difference is 29.90%.

The material was separated into three components by sublimation. The first sublimate, which was the major component, was obtained as a white solid,  $\underline{I}$ . A second higher boiling component,  $\underline{II}$ , was obtained as a brown oil in very small amounts. The third component was a brown solid residue which did not sublime even at 200°C. The infrared spectrum of  $\underline{I}$  is shown in Fig. 33.

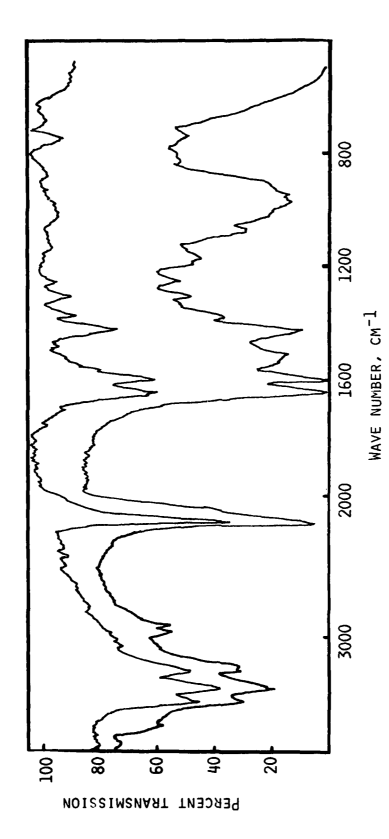
The mass spectrum of  $\underline{I}$ , shown in Fig. 34, gives a molecular weight of 123. The combined infrared and mass spectral data identify  $\underline{I}$  as 3,5-diamino-2,4-hexenenitrile, CH<sub>3</sub>-C=CH-C=N. This is consistent with the NH<sub>2</sub> NH<sub>2</sub>

fragmentation pattern tabulated in Table 10. It is likely that  $\underline{I}$  also contains some diacetonitrile, CH<sub>3</sub>-C=CH-CN,  $\underline{III}$ , (MW 82). The infrared NH<sub>2</sub>

spectra of these two compounds are indistinguishable.

Solid probe mass spectral data of the original material from the anode of cell X-8 show that there are three components which can be at least partially separated below  $250^{\rm OC}$ . A solid residue still remained after heating the sample to  $250^{\rm OC}$  in the mass spectrophotometer. The presence of  $\underline{\mathbf{I}}$  is clearly seen in the mass spectrograph of the original material shown in Fig. 35.

The other species seen in the mass spectrum of the original material were not identified. However, one component appears to have a molecular weight of 207. The infrared spectrum shows no carbonyl groups and the elemental analysis shows a C/N ratio of 2. Thus it appears that these other species are probably longer polymers of  $CH_3CN$ , possibly containing sulfur.



Infrared spectra of the product formed in the anode compartment of a  ${\rm Li/SO_2}$  cell during forced overdischarge (lower curve) and a 3,5-diamino-2,4-hexenenitrile (upper curve) after sublimation from the anode product. Fig. 33.

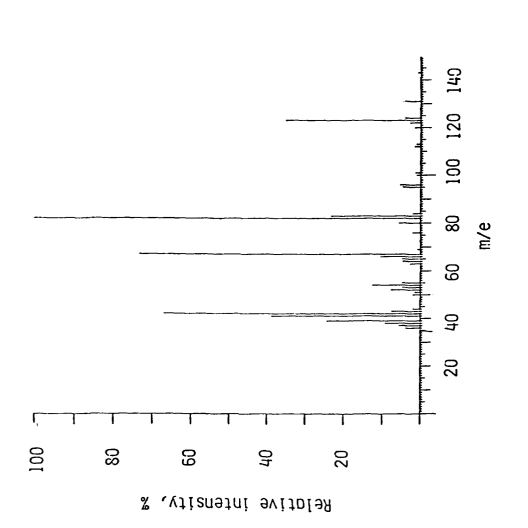
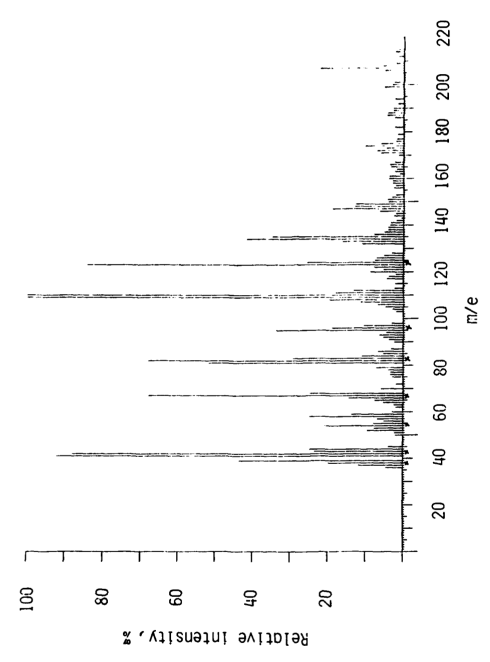


Fig. 34. Mass spectrum of 3,5-diamino-2,4-hexenenitrile,  $\underline{\mathbf{I}}$ .

TABLE 10  $$\tt MASS$  SPECTRAL DATA OF  $\underline{\tt I}$ 

| M/e | Assignment (all positive ions)   |
|-----|--|
| 123 | CH <sub>3</sub> C=CH~C=CH-CN<br>     <br>NH <sub>2</sub> NH <sub>2</sub> |
| 96  | $CH_3C=CH-C-CH$ $I$ $NH_2$ $NH_2$  |
| 82  | H <sub>2</sub> N-C=CH-C-CH<br> <br>NH <sub>2</sub>                       |
| 67  | NH <sub>2</sub> -C=CH-CN   |
| 54  | HC-C=CH<br> <br>NH <sub>2</sub>  |
| 42  | CH <sub>3</sub> CNH  |
| 41  | CH <sub>3</sub> CN   |
| 39  | C <sub>3</sub> H <sub>3</sub> or CHCN                                    |



Mass spectrum of the product isolated in the anode of cell X-8 after a discharge into reversal. \* indicates peaks from I. Fig. 35.

Infrared and qualitative data indicate that the sulfuroxy species present in the original anode product are a mixture of Li<sub>2</sub>S<sub>2</sub>O<sub>4</sub> and Li<sub>2</sub>SO<sub>3</sub>.

It is unlikely that the formation of the anode products is electrochemical in nature since the potentials of the anodes in the Type X and Type Z cells differ substantially during forced overdischarge. Experiments with a series of cells (X-18, X-19, X-20, and X-21) showed that the formation of the anode products requires a few hours of forced overdischarge (at 150 mA) and was not the result of a temperature increase at the end of discharge. Thus the products most probably result from chemical reactions between Li and acetonitrile.

Lithium and acetonitrile apparently do not react in the presence of a sufficient concentration of  $SO_2$  because of the formation of a passivating film, presumably of  $\text{Li}_2S_2O_4$ , on the Li surface (20). However, during forced overdischarge the  $SO_2$  concentration could decrease to a level where Li and acetonitrile react. Possible mechanisms for the formation of  $\underline{\text{I}}$ ,  $\underline{\text{III}}$  and  $CH_4$  are:

$$CH_3CN + 2Li \rightarrow CH_3^- + CN^- + 2Li^+$$
 (3)

$$CH_3^- + CH_3CN \rightarrow CH_4 + ^-CH_2CN$$
 (4)

$$CH_3C=N + CH_2CN \rightarrow CH_3-C-CH_2CN \qquad (5)$$

$$CH_{3}-C-CH-CN \rightarrow CH_{3}-C=CH-CN$$

$$NH H NH_{2}$$

$$III$$

$$III$$

The formation of Li<sub>2</sub>SO<sub>3</sub> in the anode compartment probably resulted from the reaction between SO<sub>2</sub> and organic Li compounds, e.g., LiCH<sub>2</sub>CN. Supporting this, we have found that reactions between  $n-C_4H_9Li$  and SO<sub>2</sub> result mainly in Li<sub>2</sub>SO<sub>3</sub>.

Other organic compounds such as 4-amino-2,6-dimethylpyrimidine (MW 123) acetoacetonitrile, 6-phenyl-2-pyridone, etc. have reportedly been identified in cells after forced overdischarge (21). We found no evidence for such species in the cells we examined. While it is possible that com-

pounds such as 4-amino-2,6-dimethylpyrimidine could form in the cell with reactions similar to those of Eqs. 3-8 (22-24), it is equally likely that some of the compounds could have formed during isolation procedures. For instance, acetoacetonitrile which was identified after extraction into a CHCl $_3$  layer from an HCl solution (21) probably resulted from the hydrolysis of I or III.

In Section 5.3.1 it was noted that a compound believed to be an aldehyde was found in two cells after forced overdischarge. This compound probably resulted from the hydrolysis of  $\underline{I}$  or  $\underline{III}$ , possibly from trace amounts of  $\underline{H}_2O$  in the two cells.

The formation of methane and other organic compounds has previously been explained by the reaction between Li (or LiAl) and acetonitrile. However, it has been assumed that the reaction occurred on the cathode where Li deposited during forced overdischarge (21). Although we found evidence for Li/Al alloy we have no evidence in the cathodes of forced overdischarged cells for unalloyed Li or any reaction products such as those found on the anode.

# 5.5 Comparison of the Behavior of Type X and Type Z Cells during Forced Overdischarge

The results of the room temperature forced overdischarge studies show that the forced overdischarge of Type Z cells presents considerable safety hazards at currents >600 mA. In actual use, it is likely that a few cells would display unsafe behavior even at lower currents; however, this would have to be verified with extensive testing of a large number of cells. The Type X Li/SO<sub>2</sub> cells appear to have a much greater resistance to forced overdischarge abuse.

A major difference in the behavior of the Type X and Type Z cells is the amount of capacity each cell yields above 0 volts.

At low discharge rates, the Type Z cell yields a slightly greater capacity to 2.0V than the Type X cell. Both types of cells yield little capacity between 2 and 0 volts at low discharge rates. At higher discharge rates (>300 mA) there are substantial differences in the capacities of the two types of cells. The difference in capacities are seen in Table 8 which list the capacities to 2.0V and 0.0V for cells discharged at high rates. For instance at 1000 mA the capacity of the Type Z cell is  $\sim$ 1200 mAh to 2.0V and ~2000 mAh to 0.0V, while the capacity of the Type X cell is 2650 mAh and 3200 mAh to 2.0V and 0.0V, respectively. At high discharge rates both types of cells yield a considerable capacity between 2 and 0 volts. However, this capacity is generally greater in the Type Z cell. Whether the high rate discharge capacity between 2-0 volts results in discharge products besides Li<sub>2</sub>S<sub>2</sub>O<sub>4</sub> is not certain and needs additional study. No evidence has been found to date for the formation of any products besides Li<sub>2</sub>S<sub>2</sub>O<sub>4</sub> and LiAl in the cathode. Probably, the low voltage discharge represents Li/Al alloy formation.

The important point is that even at high currents the capacity to 0.0V of the Type X cell is nearly equal to the capacity obtained at lower rates. In contrast, the capacity of the Type Z cell is approximately 60% (in the 1000 mA case) of the low rate capacity. This results in the presence of a large excess of unreacted Li and  $SO_2$  remaining in the Type Z cell when it goes into reversal during forced overdischarge.

The importance of excess, unreacted Li and  $\mathrm{SO}_2$  is apparent in the experiments with cells Z-9 and Z-10. The cells, discharged at low rates, had much less excess Li and  $\mathrm{SO}_2$  when forced into reversal. The cells displayed no adverse behavior even though they were forced overdischarged at high rates.

All of the cells which vented contained  $CO_2$ , COS,  $CS_2$ ,  $H_2S$ , S,  $C_2H_2$ , and  $C_2H_4$  along with  $CH_4$  and  $SO_2$ . All of the species except  $C_2H_4$  and  $C_2H_2$  are known to result from the reaction of  $SO_2$  and  $CH_4$  (25). The distribution of products depends on the reaction conditions (temperature, pressure, catalyst, etc.) The  $C_2H_2$  and  $C_2H_4$  probably result from the high temperature reaction of Li and  $CH_3CN$  via the formation of  $CH_3$ .

The above products were not identified in any cells which did not vent. Thus it appears that one of the high energy reactions which leads to excessive pressure buildup and subsequent cell venting is that between  $SO_2$  and  $CH_4$ . However, this reaction is not spontaneous and needs a high energy source of initiation.

A hypothesis for the initiation of the hazardous reactions during forced overdischarge is as follows. During a high rate discharge the carbon electrode in the cathode limited Type Z cell is rather inefficiently utilized, resulting in unused Li and SO2 in the cell. The resistivity of the cathode near the Al tab increases (possibly due to LiAl formation). The excessive heat, due to the increased resistance, initiates a violent reaction between excess Li (or Li/Al) and CH3CN and/or between SO2 and CH4. The formation of the large amounts of gaseous products along with the corresponding increase in temperature results in excessive pressure. The cell vents.

### 6.0 LOW TEMPERATURE FORCED OVERDISCHARGE EXPERIMENTS OF Li/SO2 CELLS

Our survey of the safety of Li/SO $_2$  cells indicated that low temperature forced overdischarge of Li/SO $_2$  could be particularly hazardous. Our investigation of the low temperature behavior of Li/SO $_2$  cells confirmed this. The results, discussed below indicate that forced overdischarging of Li/SO $_2$  cells at low temperature can present a serious hazard even at low currents.

Table 11 summarizes the results of forced overdischarge experiments at ~25  $^{\rm OC}.$ 

One Type X cell, X-23, was forced overdischarged at 300 mA. The voltage and temperature profiles are shown in Fig. 36. The cell capacity above zero volts was 2700 mAh. The cell voltage remained at  $\sim$  -lV for three hours. Then it began to rapidly fluctuate between -4V and -12.0V. During the forced overdischarge there were a number of small temperature increases. At about the 73rd hour, a large, sharp temperature increase occurred and the cell potential rose to just under 0 volts.

The cell did not vent. The cell was warmed to room temperature before being opened. The infrared spectrum of the gases collected from the cell after it was opened is shown in Fig. 37. Analyses of the gas mixture showed the presence of  $SO_2$ ,  $CH_4$ , COS,  $CS_2$ ,  $H_2S$  and a small amount of  $CO_2$ . Interestingly enough, no  $C_2H_4$  or  $C_2H_2$  was present.

Examination of the cell showed a series of areas where the separator was burned away. This can be seen in Fig. 38 which shows a picture of the separator and disassembled cell.

Infrared analyses of the cathode revealed that the  $\rm Li_2S_2O_4$  on the cathode had decomposed at those areas which were adjacent to the spots where the separator burned. The IR spectrum in Fig. 39 is identical to that obtained after heating a cathode sample to >170°C. All the burned areas were located in the outer two wraps of the cathode.

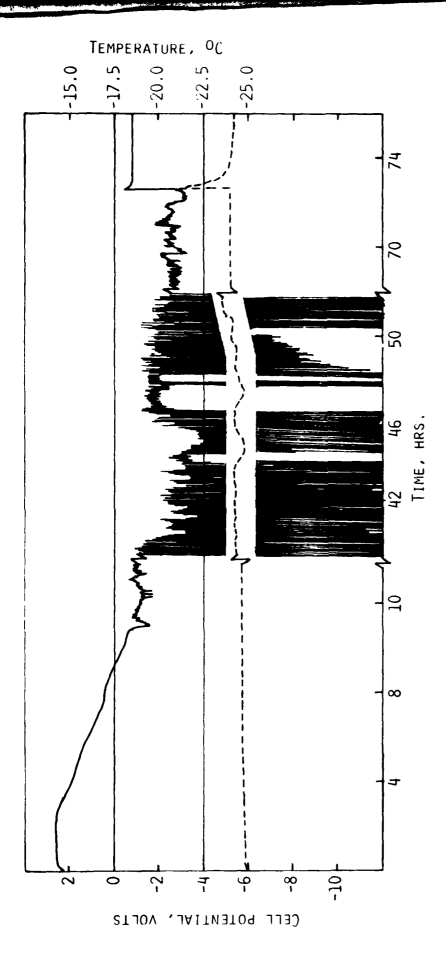
Large areas of the remainder of the cathode were covered with plated Li. These areas were very reactive. Scraping the cathode with a metal spatula resulted in small, localized explosions accompanied by an orange flame.

It was also noted that none of the brown material was present in the anode compartment.

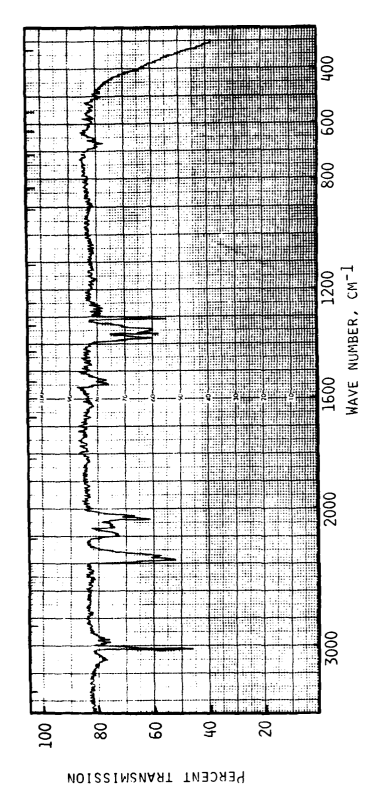
Cell X-24 was discharged and forced overdischarged under identical conditions and did not vent. Analysis of the gases in the cell showed only  $SO_2$ . While the cathode was being removed from the can (under an Ar atmosphere) it violently decomposed. The decomposition resulted in a burst of

SUMMARY OF THE RESULTS OF FORCED OVERDISCHARGE STUDIES OF Li/SO2 CELLS AT -25°C TABLE 11

| Results                                     | Li plated on cathode; areas<br>of cathode and separator<br>burned. | Cathode decomposed during removal from cell. | Vented with flame. | Vented with flame. | Did not vent. | Did not vent. |
|---|--|--|--------------------|--------------------|---------------|---------------|
| Extent of Forced<br>Overdischarge<br>(mAh)  | 19,200   | 11,850                                       | 3000               | 1500               | 3700          | 2400          |
| (mAh) to 0.0V                               | 2700   | 2700   | 1500               | 2100               | 2400          | 2000          |
| Capacity (mAh) to 2.0V 0.0V                 | 1200   | 1410   | 780                | 1200               | 1420          | 1150          |
| Forced-<br>Overdischarge<br>Current<br>(mA) | 300  | 300  | 300                | 150                | 100           | 100           |
| Discharge<br>Current<br>(mA)                | 300  | 300  | 300                | 150                | 100           | 100           |
| Ce11  | x-23   | X-24   | 2-11               | z-12               | 2-13          | 2-14          |

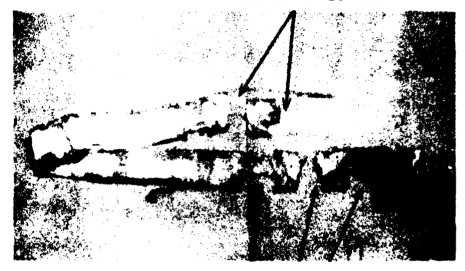


Voltage ( $\dots$ ) and temperature (---) profiles of the low temperature (-25°C) discharge and forced overdischarge of cell X-23 at 300 mÅ. Fig. 36.



Infrared spectrum of the gases formed in Li/SO2 cell X-23 during a forced overdischarge at -25°C and a current of 300 mÅ. Fig. 37.

# SEPARATOR BURNED



SEPARATOR BURNED

Fig. 38. Photograph shows the separator from cell X-23 after a forced overdischarge at -25°C. Note the burned areas which occurred near the outside of the wound electrode.

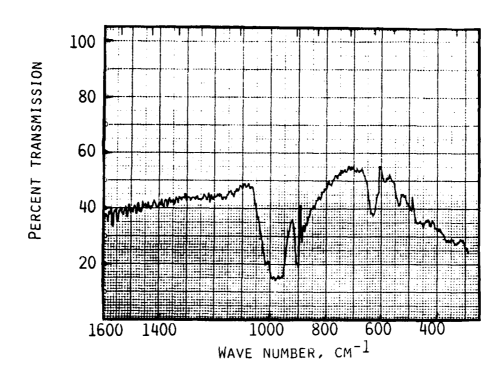


Fig. 39. Infrared spectrum of a burned area of the cathode from  $\text{Li/SO}_2$  cell X-23.

flame from the cell lasting 5-10 seconds. Subsequent analysis of the cell remains showed a complete decomposition of the  $\text{Li}_2\text{S}_2\text{O}_4$  in the cathode. The CH<sub>3</sub>CN had been removed from the cell prior to the explosion thus the reaction clearly was not that between Li and CH<sub>3</sub>CN. The reaction probably involved the highly reactive Li which plated onto the cathode during forced overdischarge, as found in cell X-23.

The results from these two cells suggest that the cathodes are in an extremely unstable state after a forced overdischarge at -25°C. Thus it is possible that a Type X cell forced overdischarged under similar conditions may explode if dropped. However, this was not verified.

Four Type Z cells were forced overdischarged at  $-25^{\circ}\text{C}$  as described in Table 11. Voltage and temperature profiles of the cells are given in Figs. 40-43. Cells Z-11 and Z-12 forced overdischarged at 300 and 150 mA, respectively, vented with flame. Cells Z-13 and Z-14 both forced overdischarged at 100 mA did not vent.

As found in the room temperature experiments, the capacities of the Type Z cells to 2.0 and 0.0 volts (see Table 11) were less than in the Type X cell. Upon reversal, the potentials of all four cells remained constant at slightly below zero volt. At this potential the most likely reaction is the plating of Li onto the cathode.

Cell Z-13 was warmed to room temperature before it was opened.  $SO_2$  was the only gas present. Unlike the Type X cells there was no plated Li visible on the cathode surface. An infrared spectrum of the cathode showed only  $\text{Li}_2S_2O_4$ . The Al grid of the cathode was fragmented and in areas completely missing. An X-ray diffraction analysis, tabulated in Table 12, showed  $\text{Li}_2S_2O_4$ , small amounts of Al and weak lines that likely correspond to LiAl. There was no evidence of unalloyed Li in the cathode.

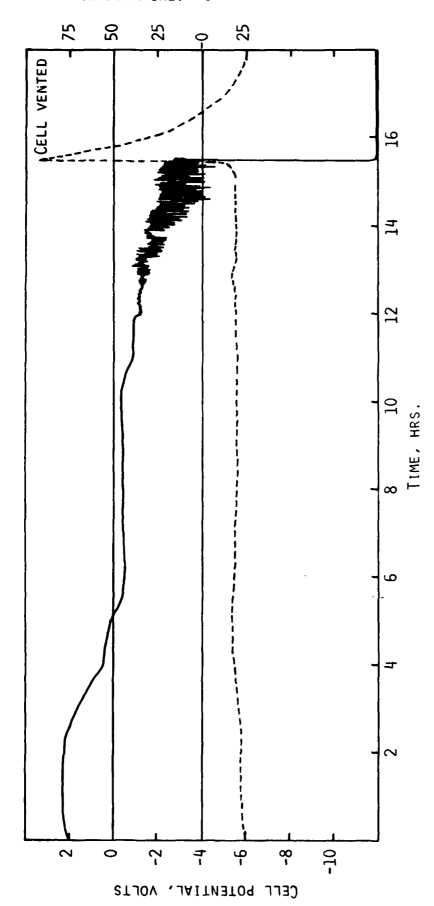
Cell Z-14 was opened immediately after the forced overdischarge at -25°C. The analysis of the cell gave identical results to that of cell Z-12, again showing no unalloyed Li on the cathode. No products, except excess Li, were found in the anode compartment of either cell.

Neither cathode displayed any evidence of shock sensitivity as found with the Type X cathodes.

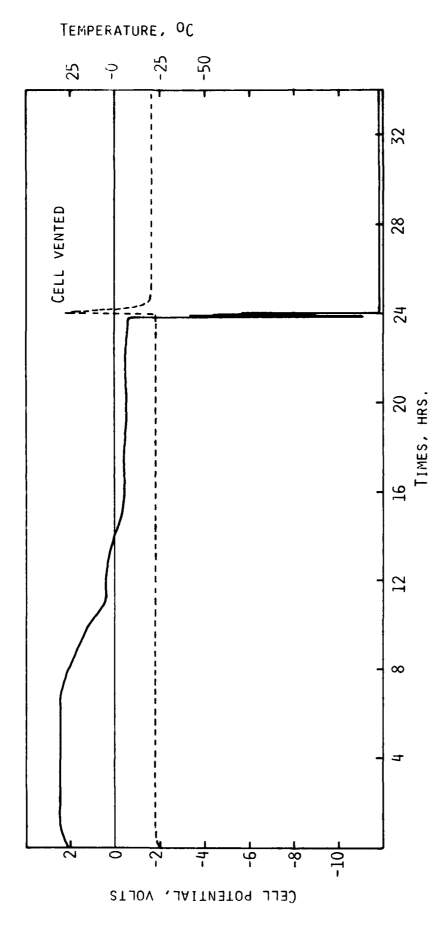
Analyses of the gases released from cells Z-11 and Z-12, confirmed the presence of COS, CS<sub>2</sub>, CH<sub>4</sub>, C<sub>2</sub>H<sub>4</sub>, CO<sub>2</sub>, H<sub>2</sub>S and SO<sub>2</sub> as previously found in the room temperature studies.

The cathodes in both cells were completely fused together. Thus the location where the explosive reaction began could not be identified. Qualitative analyses of the charred cell components confirmed the presence of S,  ${\rm SO_3}^{-2}$ , and  ${\rm SO_4}^{-2}$ .



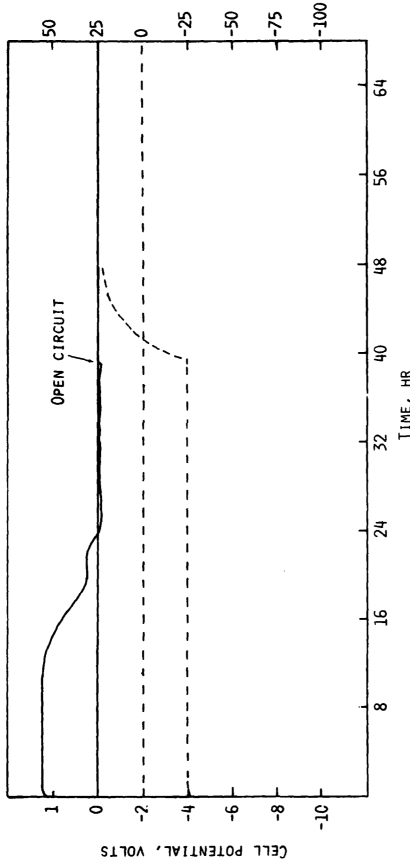


Voltage (——) and temperature (---) profiles of the low temperature (-25°C' discharge and forced overdischarge of Li/SO<sub>2</sub> cell Z-11 at 300 mA. Fig. 40.

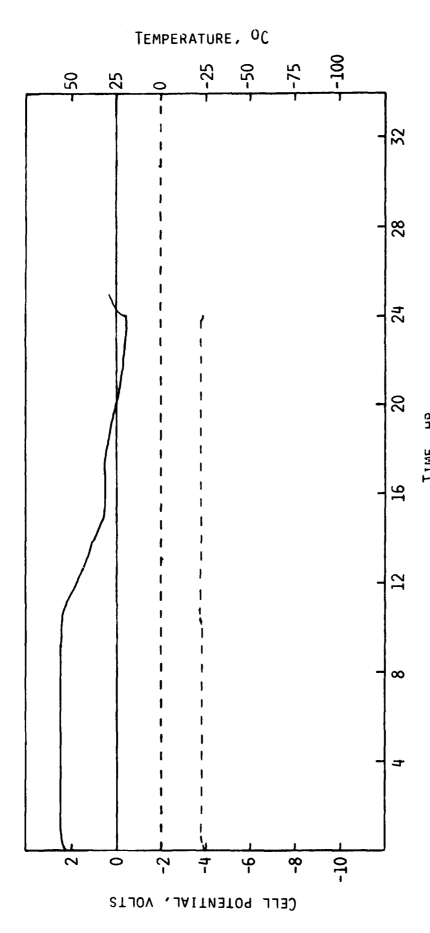


Voltage (\_\_\_) and temperature (---) profiles of the low temperature (-25 $^{\circ}$ C) discharge and forced overdischarge of Li/SO<sub>2</sub> cell 2-12 at 150 mA.





Voltage (——) and temperature (---) profiles of the low temperature (-25°C) discharge and forced overdischarge of Li/S0<sub>2</sub> cell 2-13 at 100 mA. Fig. 42.



Voltage (——) and temperature (---) profiles of the low temperature (-25°C) discharge and forced overdischarge of Li/S0 $_2$  cell 2-14 at 100 mA. Fig. 43.

TABLE 12

X-RAY<sup>a</sup> DIFFRACTION OF THE CATHODE OF CELL Z-13

| Cell        | z-13 <sup>b</sup>      | LiA  | 1                | _A.         | <u> </u>               |
|-------------|------------------------|------|------------------|-------------|------------------------|
| <u>d, A</u> | <u>I/I<sub>0</sub></u> | d, A | I/I <sub>O</sub> | <u>d, A</u> | <u>1/1<sub>0</sub></u> |
| 5.06        | 20                     |      |                  |             |                        |
| 4.33        | 80                     |      |                  |             |                        |
| 4.07        | 10                     |      |                  |             |                        |
| 3.69        | 100                    | 3.65 | 75               |             |                        |
| 3.21        | 90                     |      |                  |             |                        |
| 2.93        | 50                     |      |                  |             |                        |
| 2.74        | 15                     |      |                  |             |                        |
| 2.65        | 100                    |      |                  |             |                        |
| 2.53        | 70                     |      |                  |             |                        |
| 2.43        | 10                     |      |                  |             |                        |
| 2.33        | 20                     |      |                  | 2.34        | 100                    |
| 2.25        | 60                     | 2.26 | 100              |             |                        |
| 2.01        | 10                     |      |                  | 2.02        | 100                    |
| 1.91        | 50                     | 1.92 | 75               |             |                        |
| 1.79        | 5                      |      |                  |             |                        |
| 1.65        | 10                     |      |                  |             |                        |
| 1.61        | 10                     |      |                  |             |                        |
| 1.58        | 10                     | 1.58 | 60               |             |                        |
| 1.58        | 5                      |      |                  |             |                        |
| 1.46        | 20                     | 1.46 | 60               | 1.47        | 22                     |
| 1.41        | 10                     |      |                  |             |                        |
| 1.29        | 30                     | 1.30 | 100              |             |                        |
| 1.26        | 5                      |      |                  |             |                        |
| 1.22        | 20                     | 1.22 | 75               | 1.22        | 24                     |

 $<sup>^{</sup>a}\text{Debye-Scherrer}$  method,  $\text{CuK}_{\alpha}$  radiation.

 $<sup>^{</sup>b}$ Cell was discharged and forced overdischarged at 100 mA at -25 $^{o}$ C.

## 6.1 Discussion

The low temperature forced overdischarge of both Type X and Type 2 cells can result in hazardous cell behavior at relatively low rates.

Cell X-23 and X-24 are the only cells in which unalloyed Li was found on the carbon cathode. In some areas where the Li had plated on the cathode of cell X-23, small localized reactions occurred during forced overdischarge in which the temperature probably exceeded  $170^{\circ}$ C (the decomposition temperature of  $\text{Li}_2\text{S}_2\text{O}_4$ ).

These reactions on the cathode do not appear to be initiated by the reaction of plated Li and  $CH_3CN$ . We believe a likely alternative is the reaction between Li and  $SO_2$  ( $SO_2$  absorbed in the carbon), catalyzed by the carbon surface, i.e., a short circuited cell. The excessive heat from this reaction probably initiates reactions between Li and  $CH_3CN$ ,  $SO_2$  and  $CH_4$  and the decomposition of  $Li_2S_2O_4$ .

The fact that no unalloyed Li was found on the cathode of Type 2 cell forced overdischarged at 100 mA suggest that the exposed Al grid immediately reacts with any plated Li to form Li/Al alloy. Whether this is beneficial to the cell safety is uncertain.

The reaction in the anode compartment which produces the brown product mixture and  $CH_4$  apparently does not occur at -25 $^{O}C$  in either type of cell.

Clearly, additional work is needed in the area of low temperature use of  $\text{Li/SO}_2$  cells to evaluate their safe limits of use.

## 7.0 INVESTIGATION OF THE CHARGING BEHAVIOR OF Li/SO2 CELLS

The safety aspects of charging the  $\text{Li/SO}_2$  cell were briefly investigated with four cells. The experiments are summarized in Table 13.

Cell X-25 was discharged at 100 mA for 16 hours ( $\sim 50\%$  of the cell capacity) then recharged an equivalent amount. The voltage profile is shown in Fig. 44. The cell displayed no unsafe behavior. Post-mortem examination (2 days after the recharge) of the cell showed SO<sub>2</sub> was the only gas present. The cathode contained no detectable Li<sub>2</sub>S<sub>2</sub>O<sub>4</sub> (see Section 4.1.1). Li was the only product identified in the anode compartment. The analysis of this cell clearly shows that the cathode reaction is reversible.

Cell Z-15 was discharged and recharged under identical conditions. The voltage profile is shown in Fig. 45. The cell was not opened until two days after being charged. Post-mortem analysis of the cell showed only SO<sub>2</sub> in the gas phase. However, unlike cell X-25 the cathode contained Li<sub>2</sub>S<sub>2</sub>O<sub>4</sub>. The weight of the cathode indicated the amount of Li<sub>2</sub>S<sub>2</sub>O<sub>4</sub> was equivalent to a discharge of ~2300 mAh - more than the 1600 mAh the cell was initially discharged.

A possible reason why the Type Z cell did not recharge is that a dendritic short developed across the polypropylene separator and carried most of the charge current. Later, when the cell was stored for two days the short allowed the cell to self-discharge, thus producing additional  $\text{Li}_2\text{S}_2\text{O}_4$  in the cathode. The non-woven polypropylene separator used in the Type X cell is known to be more resistant to dendritic shorting in secondary Li cells.

Cells X-26 and Z-16 were both forced overdischarged at 100 mA before being recharged as indicated in Table 13. The voltage profiles are shown in Figs. 46 and 47. Neither cell vented.

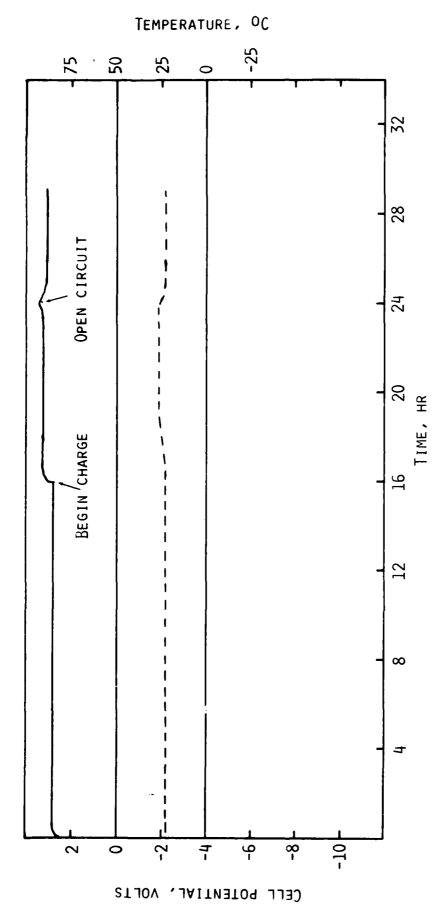
An analysis of the cells found only  $CH_4$  in gas phase. IR analyses of the cathodes of both cells showed only  $\text{Li}_2S_2O_4$ . The anode compartment of both cells contained the yellow-brown anode product typically found in forced overdischarged cells and very little Li.

Examination of cell X-26 revealed that the electrode had been misaligned during cell manufacture. The inner 3 cm of the cathode ( $\sim 10\%$  of the length) contained no Li<sub>2</sub>S<sub>2</sub>O<sub>4</sub> (via IR and qualitative tests) and had the flexible texture of a fresh cathode. The capacity to 0V (3100 mAh) was also  $\sim 10\%$  lower than obtained from other Type X cells discharged under similar conditions. This indicates that the electrode misalignment existed

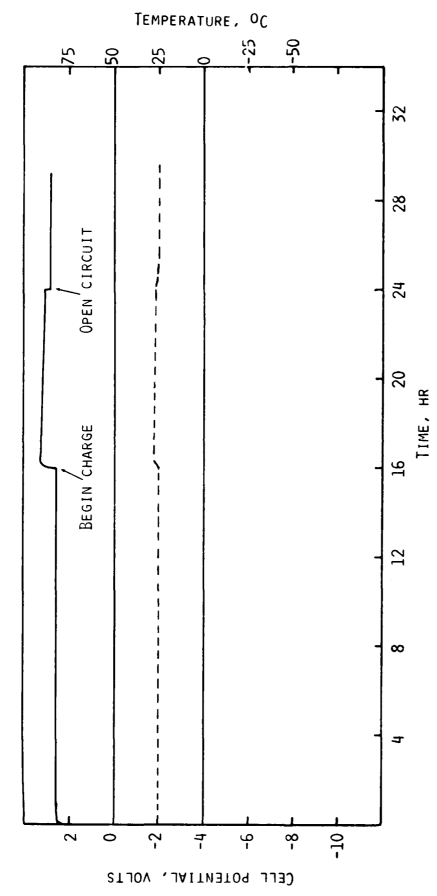
TABLE 13 SUMMARY OF THE CHARGING STUDIES OF Li/SO2 CELLS

|       |                 | Forced-         |              |                    |          |                     |                        |
|-------|-----------------|-----------------|--------------|--------------------|----------|---------------------|------------------------|
|       | Discharge       | Overdischarge   | Charge       | Discharge Capacity | Capacity | Extent of Forced    |                        |
| Ce 11 | Current<br>(mA) | Current<br>(mA) | Current (mA) | (mAh) to 2.0V      | 0.00     | Overdischarge (mAh) | Extent of Charge (mAh) |
| x-25  | 100             | ı               | 200          | 1600ª              |          | ı                   | 1600                   |
| 2-15  | 100             | 1               | 200          | 1600a              | ı        | •                   | 1600                   |
| X-26  | 100             | 100             | 100          | 3290               | 3100     | 3800                | 2500                   |
| 2-16  | 100             | 100             | 100          | 3610               | 3650     | 2850                | 5080                   |

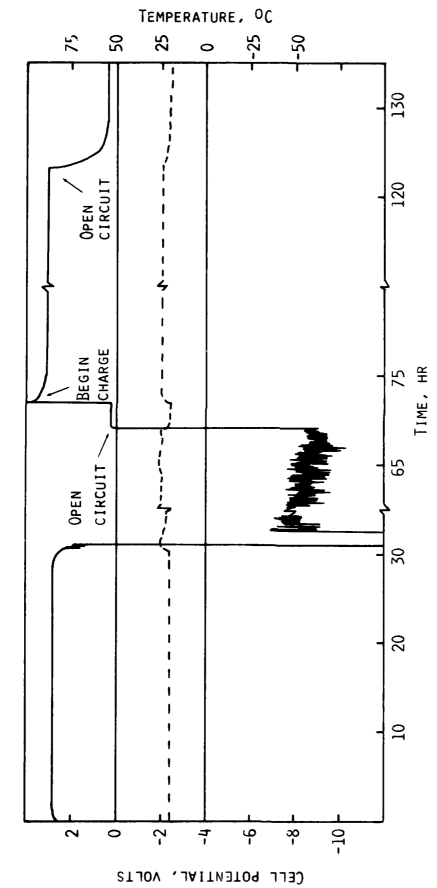
 $a\sim508$  of the normal capacity to 2.0V.



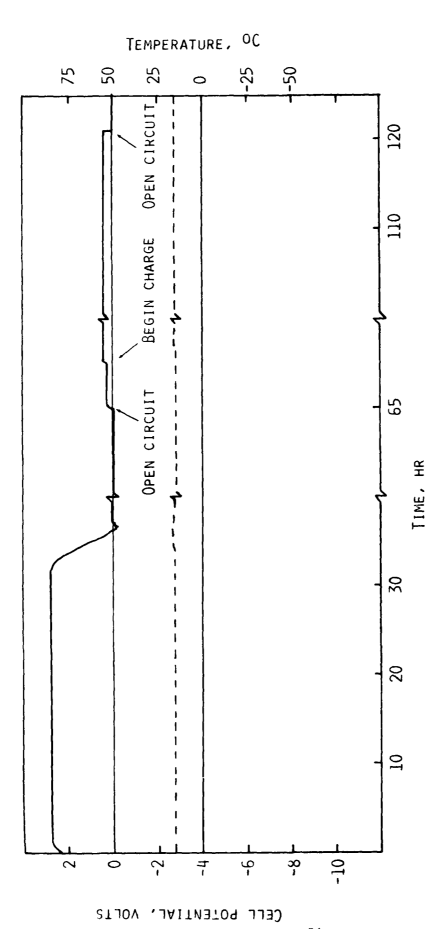
Voltage (---) and temperature (---) profiles of the discharge (100 mA) and charge (200 mA) of  $\text{Li/SO}_2$  cell X-25. Fig. 44.



Voltage (——) and temperature (---) profiles of the discharge (100 mA) and charge (200 mA) of Li/SO<sub>2</sub> cell z-15. Fig. 45.



Voltage ( $^{---}$ ) and temperature ( $^{---}$ ) profiles of the discharge, forced overdischarge, and charge of Li/SO2 cell x-26 at 100 mA. Fig. 46.



Voltage (——) and temperature (---) profiles of the discharge, forced overdischarge, and charge of Li/SO $_2$  cell 2-16 at 100 mA. Fig. 47.

in the cell before the test and was not the result of the discharge or charge process. The 3 cm of excess Li was wrapped around the outside of the electrode package and was covered by a black solid. This was not examined further.

If this cell had been used in a series connected battery it could have been forced into reversal even under non abusive discharge conditions. This example shows that the manufacturing procedures are an area that must be considered in the development of safe  $\text{Li/SO}_2$  cells.

Neither the Type X nor the Type Z cell could be recharged after forced overdischarge. The exact cause of this is not known at this time.

EIC LAGS INC NEWTON MA
INVESTIGATION OF LITHIUM SULFUR DIOXIDE (LI/SO2) BATTERY SAFETY-ETC(U)
APR 82 K M ABRAHAM, M W RUPICH, L PITTS
N60921-81-C-0084
NSWC-TR-82-148
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#### 8.0 SUMMARY

A literature and user survey of the safety hazards of Li/SO<sub>2</sub> cells and batteries identified three conditions under which the use of the system can present hazards: room temperature forced overdischarge; low temperature forced overdischarge, particularly with reversal; and, increased vulnerability to abuse (e.g., shorts, high current pulses, overdischarge or incineration) of partially discharged and stored cells.

The chemistry associated with the discharge of the Li/SO $_2$  cell was investigated in detail. An analytical procedure for the quantitative determination of Li $_2$ S $_2$ O $_4$  in discharged Li/SO $_2$  cells was developed. Quantitative determinations of Li $_2$ S $_2$ O $_4$  in several discharged cells showed very good agreement with the reaction stoichiometry of

$$2Li + 2SO_2 \rightarrow Li_2S_2O_4 \tag{9}$$

The formation of  $\rm Li_2S_2O_4$  as the principal discharge product in  $\rm Li/SO_2$  cells was further established by infrared, X-ray, and ESCA analyses of discharged cathodes.

The forced overdischarge behavior of two types of commercial C-size Li/SO $_2$  cells was investigated at 25°C. Type Z cells were found to be resistant to extended forced overdischarge at currents  $\leq 300$  mA. At currents of  $\geq 900$  mA, Type Z cells consistently vented during forced overdischarge. Type Z cells did not vent when forced overdischarged at 600 and 800 mA; however, post-test analyses of the cells revealed burned areas within the cells. Resistive heating of the Al cathode tab or areas near to it during forced overdischarge appears to initiate explosive reactions between Li and CH $_3$ CN and/or SO $_2$  and CH $_4$ . These reactions cause the cell to vent. All cells which vented produced CS $_2$ , CO $_2$ , COS, H $_2$ S, S, CH $_4$ , C $_2$ H $_4$ , C $_2$ H $_2$ , Li $_2$ S, and Li $_2$ SO $_3$ .

Type X cells were found to have good resistance towards extended forced overdischarge. No Type X cells vented when discharged and forced discharged at currents up to 1290 mA.

Post-test analyses of both Type X and Type Z cells, which did not vent during room temperature forced overdischarge, revealed methane in the gas phase and a solid product in the anode compartment. The latter was identified as a mixture of 3,5-diamino-2,4-hexenenitrile, Li<sub>2</sub>S<sub>2</sub>O<sub>4</sub>, Li<sub>2</sub>SO<sub>3</sub>, and other organic compounds. The organic compounds apparently result from reactions of excess Li in the anode compartment and CH<sub>3</sub>CN.

The forced overdischarge behavior of both Type X and Type Z cells was also investigated at  $-25^{\circ}\text{C}$ . Both types of cells were found to exhibit hazardous behavior under these conditions. The Type Z cells vented during forced overdischarge at currents of  $\geq 150$  mA. The same products, as were found in cells which vented during room temperature forced overdischarges, were found to form in cells which vented during forced overdischarge at  $-25^{\circ}\text{C}$ .

The Type X cells did not vent during forced overdischarge at  $-25^{\circ}C$ . Li was found to plate on the carbon cathode of the Type X cell during forced overdischarge at  $-25^{\circ}C$ . The cathode containing the plated Li is very reactive. Examination of one cell showed that localized, exothermic reactions occurred on the cathode surface during forced overdischarge. The reactions appear to involve plated Li and SO<sub>2</sub> absorbed in the carbon cathode and are likely catalyzed by the carbon surface.

The behavior of  $\text{Li/SO}_2$  cells during recharge was briefly investigated. Neither the Type X nor the Type Z cell displayed unsafe behavior when recharged following either a partial discharge or a forced overdischarge. The cathodic reaction was found to be reversible in Type X cells, which were only partially discharged. Type Z cells could not reversibly be recharged because of dendritic shorts across the separator.

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